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Environmental Protection DepartmentOperations and Regulatory Affairs Division

LLNL NESHAPs 2002 Annual Report



Lawrence Livermore National Laboratory

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U.S. Department of Energy Radionuclide Air Emission Annual Report (under Subpart H of 40 CFR Part 61) Calendar Year 2002

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Lawrence Livermore National Laboratory NESHAPs 2002 Annual Report

This annual report is prepared pursuant to the National Emission Standards for Hazardous Air Pollutants (NESHAPs; Title 40 Code of Federal Regulations [CFR] Part 61, Subpart H). Subpart H governs radionuclide emissions to air from Department of Energy (DOE) facilities.

SYNOPSIS

NESHAPs limits the emission of radionuclides to the ambient air from DOE facilities to levels resulting in an annual effective dose equivalent (EDE) of $10\,\mathrm{m}$ rem ($100\,\mu\mathrm{Sv}$) to any member of the public. The EDEs for the Lawrence Livermore National Laboratory (LLNL) site-wide maximally exposed members of the public from operations in 2002 are summarized here.

- Livermore site: 0.023 mrem (0.23 μ Sv) (43% from point-source emissions, 57% from diffuse-source emissions). The point-source emissions include gaseous tritium modeled as tritiated water vapor as directed by EPA Region IX; the resulting dose is used for compliance purposes.
- Site 300: 0.021 mrem (0.21 μ Sv) (85% from point-source emissions, 15% from diffuse-source emissions).

The EDEs were calculated using the EPA-approved CAP88-PC air dispersion/dose-assessment model, except for doses for three diffuse sources, which were calculated from measured concentrations and dose coefficients. Site specific meteorological data, stack flow data, and emissions estimates based on radionuclide usage inventory data or continuous stack monitoring data were the specific inputs to CAP88-PC for each modeled source.

SECTION I. Site Description

LLNL was established in 1952 to conduct nuclear weapons research and development. The Laboratory's mission is dynamic and has been broadened over the years to meet new national needs. LLNL serves as a national resource in science and engineering; its activities focus on global security, energy, global ecology, biomedicine, economic competitiveness, and science and mathematics education. LLNL comprises two sites—the main laboratory site located in Livermore, California (Livermore site), and the Experimental Test Facility (Site 300) located near Tracy, California. Figure 1 shows the locations of the sites. The University of California operates LLNL for DOE.

Livermore Site

LLNL's Livermore site occupies an area of 3.3 km² located about 60 km east of San Francisco, California, adjacent to the City of Livermore in the eastern part of Alameda County. In round numbers, 7 million people live within 80 km of the Livermore site; 76,700 of them live in the City of Livermore.

The Livermore site is located in the southeastern portion of the Livermore Valley, a topographical and structural depression oriented east-west within the Diablo Range of the California Coast Range Province. The Livermore Valley forms an irregularly shaped lowland area approximately 26 km long and an average of 11 km wide. The floor of the valley slopes from an elevation of approximately 200 m above sea level at the southwest corner.

The climate of the Livermore Valley is characterized by mild, rainy winters and warm, dry summers. The mean annual temperature is about 15°C. Temperatures typically range from –5°C during some pre-dawn hours in the winter, to 40°C on a few summer afternoons. The 2002 annual wind data for the Livermore site are displayed as a wind rose in Figure 2. Although winds are variable, the prevailing wind direction is from the southwest, especially during the summer. However, during the winter, the wind often blows from the northeast. Most precipitation occurs as rain between October and April with very little rainfall during the summer months. In 2002, the Livermore site received 271 mm of precipitation.

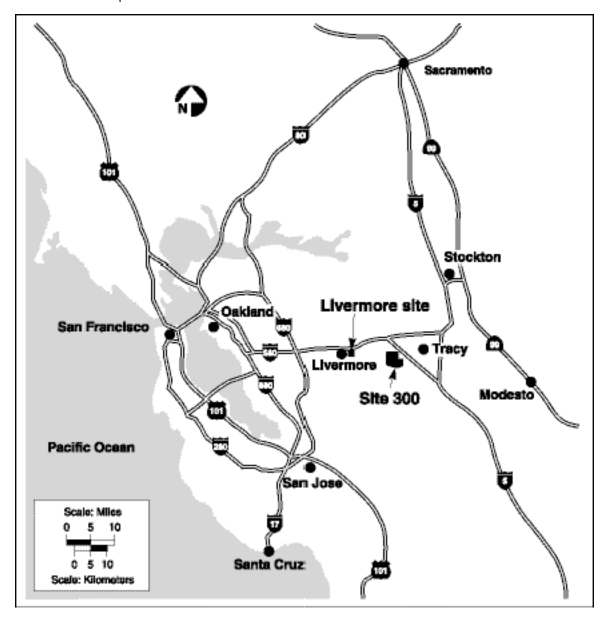


Figure 1. Locations of LLNL Livermore site and Site 300.

Site 300

Site 300, LLNL's Experimental Test Facility, is located 24 km east of the Livermore site in the Altamont Hills of the Diablo Range and occupies an area of 30.3 km². A State of California vehicular-recreation area is located nearby, and wind-turbine generators line the surrounding hills. The remainder of the surrounding area is in agricultural use, primarily pasture land for cattle and sheep. The nearest residential

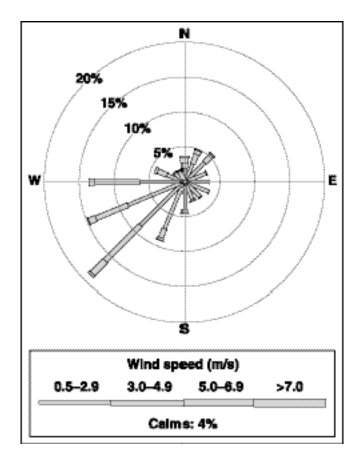


Figure 2. Wind rose showing the average annual wind speed, frequency of occurrence, and direction at the Livermore site, 2002.

area is the city of Tracy (population approximately 65,600), located 10 km to the northeast.

The topography of Site 300 is much more irregular than that of the Livermore site; it consists of a series of steep hills and ridges, which are oriented along a generally northwest/southeast trend, separated by intervening ravines. The elevation ranges from approximately 540 m in the northwestern portion of the site to 150 m at the southeast corner. The climate at Site 300 is similar to that of the Livermore site, with mild winters and dry summers. The complex topography of the site significantly influences local wind and temperature patterns, making the temperature range

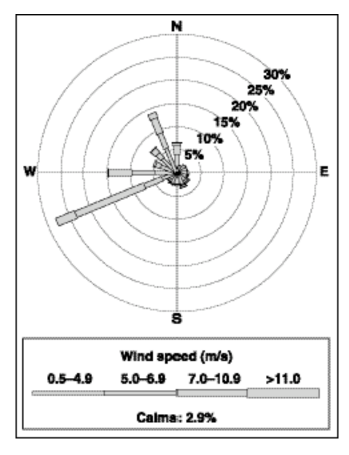


Figure 3. Wind rose showing the average annual wind speed, frequency of occurrence, and direction at Site 300, 2002.

somewhat more extreme than at the Livermore site. The 2002 annual wind data for Site 300 are displayed as a wind rose in Figure 3. Prevailing winds are from the west-southwest. As is the case at the Livermore site, precipitation is highly seasonal, with most precipitation occurring between October and April. Site 300 received 220 mm of precipitation during 2002. The mean annual temperature is about 17°C.

SECTION II. Air Emission Sources and Data

Sources

Nearly a hundred different radioisotopes are used at LLNL for research purposes, including biomedical tracers, tritium, mixed fission products, transuranic isotopes, and others—see Table 1 for a list of the radionuclides and the "radionuclides" column in the Attachment 1 spreadsheet for a breakdown by facility. Radioisotope handling procedures and work enclosures are determined for each project, depending on the isotopes, the quantities being used, and the types of operations being performed. Work places include glove boxes, exhaust hoods, and laboratory bench tops. Exhaust paths to the atmosphere range from triple HEPA (High Efficiency Particulate Air) filtered ventilation systems, to roof vents and stacks lacking abatement devices, to direct dispersal of depleted uranium during explosives testing at Site 300, to a variety of diffuse area sources.

Table 1. Radionuclides used at LLNL during 2002.

^{3}H	$^{54}\mathrm{Mn}$	⁹⁹ Tc	¹⁴⁸ Gd	²²⁹ Th	240 Pu
$^{7}\mathrm{Be}$	⁵⁵ Fe	103 Rh	151 _{Pm}	^{230}Th	²⁴¹ Am
$^{10}\mathrm{Be}$	57 _{Co}	106 Ru	151 _{Sm}	^{231}Pa	241 Pu
^{13}N	58Co	¹⁰⁹ Cd	152 _{Eu}	²³² Th	²⁴² Cm
14 _C	⁵⁹ N i	113 Sn	154 Eu	232 _U	²⁴² Pu
15 _O	⁶⁰ Co	125_{I}	155 _{Eu}	233 _U	²⁴³ Am
^{22}Na	$63_{ m Ni}$	125 _{Sb}	$^{172}\mathrm{Hf}$	234U	²⁴⁴ Cm
32 _P	75 _{Se}	$131_{ m I}$	174 Lu	235U	244 Pu
33 _P	⁸⁵ Sr	133 B a	¹⁹⁵ Au	236 _{Pu}	²⁴⁶ Cm
35 _S	88 _Y	134_{Cs}	195mPt	236 _U	²⁴⁸ Cm
36Cl	$^{90}\mathrm{Sr}$	137 _{Cs}	$^{207}\mathrm{B}\mathrm{i}$	^{237}Np	²⁴⁹ Cf
40 K	90γ	$^{140}\mathrm{B}\mathrm{a}$	^{209}Po	237 _U	²⁵⁰ Cf
41Ar	⁹⁴ Nb	141 Ce	210_{Pb}	238 Pu	²⁵² Cf
⁴¹ Ca	⁹⁵ Nb	144 Ce	223 Ra	238U	
46Sc	^{95}Zr	¹⁴⁷ Nd	226 Ra	^{239}Np	
51 _{Cr}	⁹⁹ Mo	¹⁴⁷ Pm	²²⁸ Th	²³⁹ Pu	

Sources of radioactive material emissions to air at LLNL are divided into two categories for purposes of evaluating NESHAPs compliance: point sources (including stacks, roof vents, and explosive experiments conducted on Site 300's firing tables) and diffuse area sources (including dedicated waste accumulation areas and other areas of known contamination). Several emission sources are treated as diffuse extended area sources, including Radioactive and Hazardous Waste Management's "Tank Farm" operations at Building 514 and waste storage at the Building 612 Yard, and other Livermore-site sources external to buildings. Detailed

information is given in Attachment 1 for emissions from LLNL's radiological operations that took place during 2002.

2002 Air Monitoring

In this section we describe continuous stack-effluent sampling systems at selected LLNL facilities and ambient air monitors in place at numerous locations on and off LLNL sites.

Continuous Stack Air Effluent Monitoring

Actual measurements of radioactivity in air and effluent flow are the basis for reported emissions from continuously monitored sources. In 2002, there were seven buildings (Buildings 175, 177, 235, 251, 331, 332, and 491) at the Livermore site and one building (Building 801A) at Site 300 that had radionuclide air effluent monitoring systems. These buildings are listed in Table 2, along with the number of samplers, the types of samplers, and the analytes of interest. Many samplers would operate from emergency power systems if normal power were lost.

Air samples for particulate emissions are extracted downstream of HEPA filters and prior to the discharge point to the atmosphere. Particles are collected on membrane filters. The sample filters are removed and analyzed for gross alpha and beta activity on a weekly or bi-weekly frequency depending on the facility. In most cases, simple filter aerosol collection systems are used. However, in some facilities, alpha continuous air monitors (CAMs) are used for sampling. In addition to collecting a sample of particles, the CAM units provide an alarm capability for the facility in the event of an unplanned release of alpha activity.

Detection of gross alpha and beta activity resulting from particles collected on the air filters is accomplished using gas flow proportional counters. Analysis is delayed for at least four days from the end of sample collection to allow for the decay of naturally occurring radon daughters. For verification of the operation of the counting system, calibration sources, as well as background samples, are intermixed with the sample filters for analysis. Analysis is performed by the Radiological Measurements Laboratory (RML) in LLNL's Hazards Control Department (HCD).

Each stack of the Tritium Facility (Building 331) is monitored for tritium release by both an alarmed continuous monitoring system and by molecular sieve continuous samplers. The alarmed monitors, which are Overhoff ion chambers, provide real time tritium concentration release levels (HT, HTO, or other gaseous forms). The sieve samplers discriminate between tritiated water (HTO) vapor and molecular tritium (HT); they provide the values used for environmental reporting and are exchanged weekly. Each sieve sampler (not alarmed) is in parallel with an alarmed monitor and consists of two molecular sieves. The first sieve collects tritiated water vapor; the second sieve contains a palladium-coated catalyst that converts molecular tritium to tritiated water, which is then collected. The molecular sieve samples are submitted to the Hazards Control Analytical Laboratory where they are put into a

recovery system for the bake out of tritiated water vapor and subsequent condensation and collection of the water. The retrieved tritiated water is analyzed by RML using liquid scintillation counting techniques.

Environmental Protection Department (EPD) environmental analysts review data from air particulate sampling filters and molecular sieves.

Table 2. Air effluent sampling systems and locations.

Building	Facility	Analytes	Sample type	Number of samplers
175	MARS ^a	Gross □, □ on particles	Filter	6
177	Extractor Test ^a	Gross □, □ on particles	Filter	1
235	Chemistry and Materials Science	Gross □, □ on particles	Filter	1
251	Heavy Elements Unhardened area Hardened area	Gross □, □ on particles Gross □, □ on particles	Filters Filters	24 4
331	Tritium	Tritium	Ionization Chamber ^b	4
		Gaseous tritium/ tritiated water vapor	Molecular sieve	s 4
332	Plutonium	Gross □, □ on particles	CAMb	12
		Gross □, □ on particles	Filters	16
491	Isotope Separation ^a	Gross □, □ on particles	Filters	1
801A	Contained Firing Faciltiy	Gross □, □ on particles	Filters	1

Note: "CAM" denotes Eberline continuous air monitors.

Results of Stack Monitoring for Tritium: Operations in the Tritium Facility (Building 331) in 2002 released a total of 36 Ci $(1.3 \square 10^{12} \text{ Bq})$ of tritium. Of this, approximately 33 Ci $(1.2 \square 10^{12} \text{ Bq})$ were released as tritiated water (HTO). The remaining 9.7% of the tritium released, 3.5 Ci $(1.3 \square 10^{11} \text{ Bq})$, was elemental tritium

^a Operations discontinued, however, air effluent sampling systems at this building continue to operate as part of the maintenance and surveillance shutdown plan for the facilities. The Building 177 effluent sampling system was removed in Feb. 2002, after decontamination and decommissioning of the facility was completed.

b Alarmed systems.

gas (HT). The highest single weekly stack emission from the facility was 3.8 Ci (1.4 \square 10¹¹ Bq), of which more than 99% was HTO.

Building 331 tritium emissions, as measured by stack monitoring, remained considerably lower than levels that occurred during the 1980s. We anticipate that emissions over the next five years will exceed the 2000–2002 levels, as research and development work is performed for new programmatic efforts. However, engineered controls designed to contain and recapture tritium leakage should maintain relatively low emissions. Figure 4 illustrates the combined HTO and HT emissions from the facility since 1981.

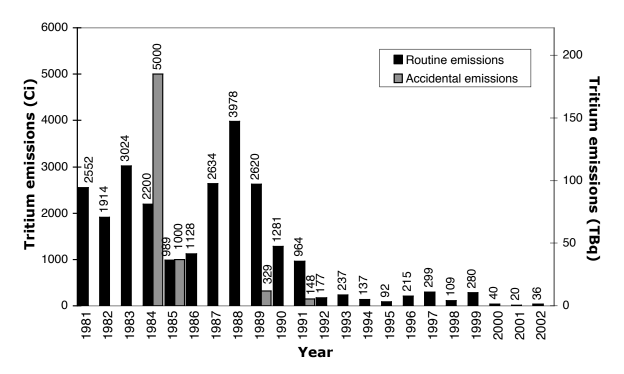


Figure 4. Combined HT and HTO emissions from the Tritium Facility, 1981–2002, distinguishing between chronic releases during normal operations (black bars) and acute accidental releases (gray bars). Accidental releases are predominantly HT gas.

Stack Monitoring for Gross Alpha and Gross Beta Radiation: For most discharge points at the other facilities where continuous stack sampling is performed, the results are below the minimum detectable concentration (MDC) of the analysis; sometimes as few as 1 to 4 samples (out of 25 to 50 per year) have concentrations greater than the MDC. Generally, these few samples having results above the MDC are only marginally above it. Use of zero values for this type of data can be justified based on knowledge of the facility, the use of tested, multiple stage, HEPA filters in all significant release pathways, and alpha spectroscopy based

isotopic analyses of selected air sampling filters. These isotopic analyses demonstrate that detected activity on air sampling filters comes from naturally occurring radionuclides, such as radon daughters, e.g., polonium, on the air sampling filters. In addition, because of exhaust configurations at some facilities, the monitoring systems sometimes sample air from the ambient atmosphere along with the HEPA filtered air from facility operations, giving rise to background atmospheric radioactivity being collected. Because of these considerations, the emissions from such facility operations are reported as zero. As a result, there are no dose consequences, and doses reported for these operations are zero. Furthermore, even if the MDC values were used in calculations of the emission estimates for these facilities, which would be an extremely conservative approach, the total dose attributable to LLNL activities would not be significantly affected.

An effluent sampling system was installed in Building 801 at Site 300 in early 2002. Although all facility operations are HEPA filtered, this building has a large high bay room that exhausts to the stack without HEPA filtration. Consequently, some of the air sampled by the effluent sampling system is essentially outside, ambient air. In order to determine if any releases actually occurred from this facility, the sampling results must be compared to ambient air. In 2002, five samples out of 37 had concentrations greater than the MDC. The median concentration of the Building 801 stack detections, $1.3 \,\Box 10^{-4} \,\mathrm{Bq/m^3}$ (3.6 $\Box 10^{-15} \,\mathrm{Ci/m^3}$), was lower than the median concentrations of the detections from two offsite sampling locations that are used to establish background levels of gross alpha and beta activity for direct comparison to results from the air effluent samplers. The median of all 37 of the five Building 801 samples, $3.0 \,\Box 10^{-5} \,\mathrm{Bq/m^3}$ (8.0 $\Box 10^{-16} \,\mathrm{Ci/m^3}$), was approximately three times lower than the median of all of the offsite sampling location samples. Therefore, it is reasonable to conclude that Building 801 operations did not have radioactive emissions.

None of the facilities monitored for gross alpha and beta had emissions in 2002.

Air Surveillance Monitoring for Radioactive Particles and Gases

Surveillance air monitoring for tritium and radioactive particles has been in place since the 1970s. LLNL currently maintains seven continuously operating, high volume, air particulate samplers on the Livermore site, nine in the Livermore Valley, eight at Site 300, and one in Tracy. LLNL also maintains twelve continuously operating tritiated water vapor samplers on the Livermore site, six samplers in the Livermore Valley and one at Site 300. The samplers are positioned to ensure reasonable probability that any significant airborne concentration of particulate or tritiated water vapor effluents resulting from LLNL operations will be detected. Many of the surveillance air monitors are placed near diffuse emission sources, such as those near Buildings 292, 331, 514, and 612, as well as in and around the Southeast Quadrant of the Livermore site. As such, their results can be used to estimate and/or confirm the emissions from the associated diffuse sources. Also included is

an air particulate monitor positioned at the location of the hypothetical maximally-exposed member of the public (defined in Section III) for the Livermore site. Data from air surveillance monitors provide a valuable test of predictions based on air dispersion modeling, and can help characterize unplanned releases of radioactive material.

The data from the surveillance air monitoring network provide continuous measurements of the concentrations of radionuclides present in the air at the Livermore site, Site 300, and in the surrounding areas. Data from the network are presented in the LLNL Site Annual Environmental Report (SAER), which is available to the public in hardcopy form, on CD, and on the Internet. (See, e.g., Gallegos et al., *Environmental Report 2001*, Lawrence Livermore National Laboratory, Livermore, CA, UCRL-50027-01, September 2002; http://www.llnl.gov/saer).

Radionuclide Usage Inventory Update

A "partial" accounting of LLNL's radiological emission sources was made in 2002 (as was done in 2001), in accordance with the allowance by EPA that a 100% accounting need be made only every third year. A 100% accounting was made when reviewing and reporting on operations conducted in 2000.

The partial accounting focused on sources in four categories: (1) the group of sources that collectively (in a ranked list) accounted for at least 90% of the dose to the maximally-exposed public individual from both the Livermore site and Site 300 in the previous year's (2001) assessment; (2) all "new" sources, i.e., those that commenced emissions in 2002, or sources that showed significantly elevated releases over 2001 levels; (3) all monitored sources; and (4) all sources in the major LLNL waste stream dealt with by Radioactive and Hazardous Waste Management (RHWM) Division in the Environmental Protection Department (EPD) of LLNL.

Radionuclide usage inventory forms, with guidance for completing them, were sent to all assurance managers, facility managers, and project-responsible persons connected with activities meeting these criteria for our partial accounting. The forms were completed by experimenters, and certified by facility managers. In particular, radionuclide usage data for all Site 300 explosives experiments and all significant stack and diffuse sources at both sites were included in this update.

SECTION III. Dose Assessment Methods & Concepts

Description of the Air Dispersion and Dose Model

Most estimates of individual and collective radiological doses to the public from LLNL operations were obtained using the EPA-developed computer code CAP88-PC. The four principal pathways—internal exposures from inhalation of air, ingestion of foodstuff and drinking water, external exposures through irradiation from contaminated ground, and immersion in contaminated air—are evaluated by CAP88-PC. The doses are expressed as whole-body effective dose equivalents (EDEs), in units of mrem/y (1mrem = 10 μ Sv). Separate doses for Livermore site and Site 300 emissions are reported. An LLNL-modified version of CAP88-PC (designated CAP88-PC-T) that contains an improved tritium model NEWTRIT (not yet approved by EPA for use in regulatory compliance evaluations), was also used in the assessment of inhalation and ingestion doses from tritium, for purposes of comparison.

Three potential doses are emphasized: (1) The dose to the site-wide maximally exposed individual (SW-MEI), which combines the contributions of all emission points to dose at a publicly-accessible facility (e.g., a business, church, school, or residence), for comparison to the 10 mrem/y (100 \$\pi\$Sv/y)\$ standard; (2) The maximum dose to any member of the public (assumed to be at the LLNL fence line), in any direction, due to each unabated emission point on the site to determine the need for continuous monitoring; and (3) the collective dose to populations residing within 80 km of the two LLNL sites, summing the products of individual doses received and number of people receiving them.

Summary of Model Input Parameters

General Model Inputs: Attachment 1 details the key identifiers and input parameters for the CAP88-PC model runs. These include building number; stack ID; isotope(s); emission rate in curies per year (1 Ci = $3.7 \square 10^{10}$ Bq); and stack parameters, including height, diameter, and emission velocity.

Meteorological Data: All model runs used actual 2002 Livermore-site and Site 300 meteorological data, collected from the meteorological towers for each site. At these towers, wind speed and direction are sampled every few seconds, temperature sampled every minute, and all are averaged into quarter-hour increments, time tagged, and computer recorded. The data are converted into a CAP88-PC input wind file using EPA guidelines.

Surrogate Radionuclides: CAP88-PC contains a library of 265 radionuclides; however, it does not contain all the radionuclides in use at LLNL. As a consequence, it was necessary in a few cases to use surrogate radionuclides to estimate EDEs. Attachment 2 shows the surrogate radionuclides used in CAP88-PC. The selection of a suitable surrogate is based upon several criteria, including metabolically similar

behavior and similar modes of decay and decay energies of the radiation type of the isotope of interest. Once a surrogate is selected, the equivalent source term is adjusted by the product of the initial inventory of the isotope of interest and the ratio of the effective dose equivalent of the surrogate to that of the isotope of interest. In some cases, experimenters did not provide isotopic analyses of mixtures of radionuclides, and they identified the radionuclides used as "gross alpha," "gross beta," "gross gamma," or "mixed fission products" (MFP). In these cases, ²³⁹Pu was used as the surrogate for gross alpha, ¹³⁷Cs was used as the surrogate for gross gamma, and ⁹⁰Sr was used as the surrogate for gross beta and mixed fission products to provide conservative dose estimates.

Population Inputs: Population distributions centered on the two LLNL sites were compiled from the LandScan Global Population 1998 Database developed by Dr. Jerome Dobson at Oak Ridge National Laboratory. The population data files (distribution of population with distance and direction) used in the 2002 modeling effort are the same as those described in the 2000 NESHAPs annual report (*LLNL NESHAPs 2000 Annual Report*, Gallegos et al., June 2001).

Land Use and Agricultural Inputs: Options for model inputs regarding agricultural characteristics and land use are established by the EPA, and the particular designation selected can strongly influence the ingestion dose received by the population being evaluated. The "user entered" option was again selected for the CAP88-PC modeling effort for 2002. The values entered corresponded to the "local agriculture" option (everything is home produced), with one exception—all milk consumed was assumed to be imported when assessing dose to individuals (as opposed to populations). An assumption that all milk comes from local cows would not be supported by the agricultural activities conducted in the area. For population dose assessments, all food is considered to be locally grown, i.e., grown within an 80 km radius about the site; default densities of agricultural products in California are used.

Emission Source Terms: The source term for each emission point in the calculations was determined by one of two methods: For continuously monitored sources, the sampling data (curies released per unit time) for each radionuclide were used directly. For unmonitored facilities, the radionuclide usage inventories, together with time factors and EPA-specified physical state factors, are used to estimate potential emissions to air from a source. Time factors are used to adjust for the fact that a radionuclide may not always be in the same facility all year or may be encapsulated or enclosed for a substantial part of the year. Time factors are chosen to allow a more reasonable estimate of the amount of radioactive material released into the atmosphere. The EPA-specified factors for potential release to air of materials in different physical states (solid, liquid, powder, or gas) are those stated in 40 CFR Part 61, Appendix D. If the material was an unconfined gas, or any material heated above 100°C (with exceptions noted in Table 3), then the factor 1.0 was used;

for liquids and powders, $1.0 \square 10^{-3}$ was used; and for solids, $1.0 \square 10^{-6}$ was used. The U.S. EPA has granted approval for LLNL to use alternative physical state factors for elemental uranium, uranium/niobium alloy, and elemental plutonium. Table 3 provides the approved temperatures for application of the physical state factor for each material.

These factors are allowed provided that the material is not intentionally dispersed to the environment and that the processes do not alter its chemical form. The physical-state-dependent release fraction and the time factor are used to adjust (by multiplication) the total annual usage inventory to yield the potential annual release to air. In addition, emission control abatement factors (40 CFR 61, Appendix D), when applicable, were applied. Each HEPA filter stage was given a 0.01 abatement factor. (However, abatement factors were not used to evaluate compliance with the 0.1 mrem [1 μ Sv] standard that determines the need for continuous monitoring at a facility.) The use of actual stack effluent sampling data is much more direct, and presumably more accurate, than using assumptions based on usage inventory, time factors, release fractions, and emission control factors.

Table 3. List of materials exempted from the "treat as a gas above 100°C rule," and temperatures at which the various physical state factors apply.

Material	Solid physical state factor	Liquid physical state factor	Gas Physical state factor	Year Approved
Elemental uranium	<1100°C	Between 1100°C and 3000°C	>3000°C	1996
Uranium/niobium al	lloy<1000°C	Between 1100°C and 3000°C	>3000°C	2001
Elemental plutonium	<600°	Between 600°C and 3000°C	>3000°C	2001

Site-Wide Maximally Exposed Individual: For LLNL to comply with the NESHAPs regulations, the LLNL site-wide maximally exposed individual cannot receive an EDE greater than 10 mrem/y $(100 \,\mu\text{Sv/y})$. The site-wide maximally exposed individual (SW-MEI) is defined as the *hypothetical* member of the public at a single residence, school, business, church, or other such facility, who receives the greatest LLNL induced EDE from the combination of all radionuclide source emissions, as determined by modeling.

At the Livermore site, the SW-MEI for 2002 was found, as usual, to be located at the UNCLE Credit Union, about 10 m outside the controlled eastern fence line of the site, but about 10 m within the perimeter of the site property, as shown in Figure 5. At Site 300, the 2002 SW-MEI was again, as in the previous two years, located at the boundary with the Carnegie State Vehicle Recreation Area, managed by the California Department of Parks and Recreation, approximately 3.2 km south southeast of the firing table at Building 851, as shown in Figure 6.

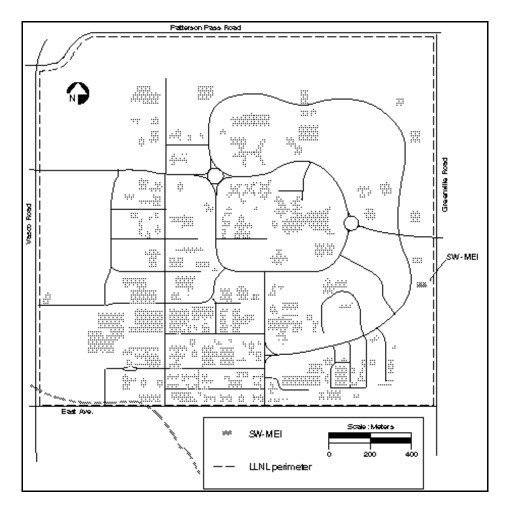


Figure 5. Location of Site-Wide Maximally Exposed Individual (SW-MEI) at the Livermore site, 2002.

In the Attachment 1 spreadsheet, the distance and direction to the respective SW-MEI are shown for each facility at each site. Doses to the SW-MEIs were evaluated for each source and then totaled for site-specific evaluations against the 10 mrem/y (100 μ Sv) dose standard (see "Total Dose to Site-Wide Maximally Exposed Individuals" in Section IV).

Maximally Exposed Public Individual: To assess compliance with the EPA requirement for continuous monitoring of a release point (potential dose greater than $0.1 \, \text{mrem/y} \, [1.0 \, \mu \text{Sv/y}]$), emissions must be individually evaluated from each point source; the location of the maximally exposed public individual (MEI) is generally different for each emission point. The maximum dose at a location of unrestricted public access typically occurs at a point on the site perimeter. Therefore, it is often referred to as the maximum "fence line" dose, although the off-site maximum dose could occur some distance beyond the perimeter. (This could

happen, e.g., when a stack is close to the perimeter; however, for all emission points at the Livermore site and Site 300, calculations show that ground level concentrations of radionuclides generally decline continuously beyond LLNL boundaries.) As stipulated by the regulations in 40 CFR Section 61.93 (b)(4)(ii), modeling for assessment of continuous monitoring requirements assumed unabated emissions (i.e., no credit was taken for emission abatement devices, such as filters). The Attachment 1 spreadsheet provides the dose to the MEI and the distance and direction to the LLNL fence line where the MEI is located.

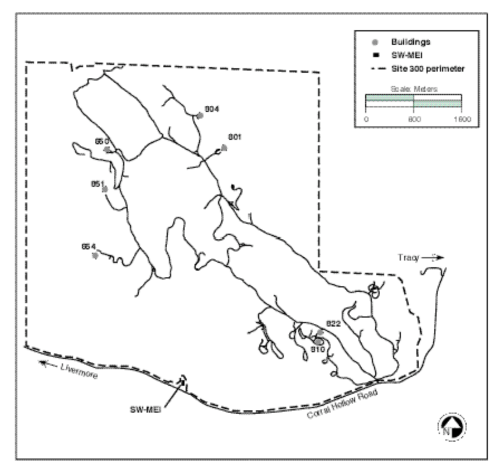


Figure 6. Location of Site-Wide Maximally Exposed Individual (SW-MEI) at Site 300, 2002.

Special Modeling Challenges

Among the sources at LLNL, explosives tests using depleted uranium at Site 300 and diffuse sources at both sites required special attention.

Site 300 Explosives Experiments: Some of the assemblies for Site 300 explosives experiments contain depleted uranium (DU) and possibly other

radioactive materials. (The radioactive material does not contribute to the explosive energy, which is entirely chemical in origin.) The explosives assemblies are placed on an open-air firing table and detonated. Only limited data are available to characterize the initial state of the cloud of explosive decomposition products created by the detonation because properties of the cloud are not routinely measured in the experiments. Empirical scaling laws can be used, however, to define the size and height of the cloud using explosives inventories. When the assembly contains DU, the three uranium isotopes with atomic weights 238, 235, and 234 are assumed to occur in the cloud in the weight percentages 99.8, 0.2, and $5 \square 10^{-4}$. Their masses are multiplied by their specific activities to determine the total activity for each isotope in the cloud. For simplicity, it is assumed that 100% of the uranium is aerosolized and dispersed as a gaseous cloud, and that the median particle size is the CAP88-PC default value of 1 μ m. These assumptions result in a highly conservative off-site dose estimation. We believe the percentage of uranium dispersed is well below the assumed value of unity probably no greater than 0.2 — but considerable time and expense would be required to justify use of a lower value. Furthermore, CAP88-PC does not capture the short-duration, explosive nature of the release; rather it simulates each shot as a low level, steady state, stack-type emission occurring over one year. An alternative modeling methodology for treating these detonations, based on a "puff" code, was submitted to EPA for approval in 1992, but LLNL was directed to use the CAP88-PC code.

Diffuse Sources: Diffuse emissions generally arise from extended-area sources external to buildings. Such sources are difficult to quantify. At present there are no EPA-mandated methods for estimation or measurement of diffuse sources; dose calculations associated with this type of source are left to the discretion of the DOE contractor facility. Dose assessments for Livermore-site and Site 300 diffuse sources are variously derived based on radionuclide usage inventory data, environmental surveillance monitoring data, samples of contaminated materials, and other methods. The doses from principal diffuse sources in 2002 are described below in Section VIII.

Modeling Documentation

Dose assessment modeling runs were conducted for all sources (point and diffuse) meeting the criteria of the reduced accounting for 2002. The model used was EPA's CAP88-PC code (see Section III). Files were incorporated for meteorological data (wind, precipitation, and temperature) and population data representing both sites, along with the 2002 radionuclide usage inventory or stack effluent monitoring data. Annual dose is reported as whole-body EDE expressed in units of mrem (followed by μ Sv; 1 mrem = 10 μ Sv). Copies of individual model runs, including input parameters and resultant calculated doses, are on file with the Terrestrial & Atmospheric Monitoring & Modeling Group (TAMM) of the Environmental Protection Department at LLNL.

SECTION IV. Results of 2002 Radiological Dose Assessment

This section summarizes the doses to the most exposed public individuals from LLNL operations in 2002, shows the temporal trends and comparison to previous years, presents the potential doses to the populations residing within 80 km of either the Livermore site or Site 300, and summarizes LLNL's compliance with 40 CFR 61, Subpart H (61.93).

Total Dose to Site-Wide Maximally Exposed Individuals

The total dose to the Livermore site SW-MEI from operations in 2002 was 0.023 mrem (0.23 μ Sv). Of this, 0.010 mrem (0.10 μ Sv) or 43% was contributed by point sources, while diffuse emissions accounted for 0.013 mrem (0.13 μ Sv) or 57% of the total. The point source dose includes Tritium Facility HT emissions modeled as HTO, as directed by EPA Region IX. The SW-MEI dose calculated using CAP88-PCT with its NEWTRIT model (see "Modeling Dose from Tritium" in Section VII), rather than the default CAP88-PC code, reduced the tritium component of the Livermore site dose from 0.020 mrem (0.20 μ Sv) to 0.015 mrem (0.15 μ Sv).

The total dose to the Site 300 SW-MEI from operations in 2002 was 0.021 $\bar{\mu}$ Sv). Point source emissions from firing table explosives experiments accounted for 0.018 mrem (0.18 μ Sv), or 85%, of this total, while 0.0033 $\bar{\mu}$ Sv), or about 15%, was contributed by diffuse sources.

Table 4 shows the facilities or sources that accounted for more than 90% of the doses to the SW-MEI for the Livermore site and Site 300 in 2002. Although LLNL has nearly 200 sources with potential for releasing radioactive material to air according to NESHAPs prescriptions, most are very minor. Nearly the entire radiological dose to the public from LLNL operations comes from no more than a dozen sources. In 2002, a proposal was made to EPA for permission to use surveillance monitoring in place of inventory-based modeling to account for dose contributions from minor sources. This proposal was approved in April 2003 (see Attachment 3), and will be implemented for next year's NESHAPs annual report.

Table 5 compares 2002 doses with those of previous years. No diffuse emissions were reported at Site 300 for years before 1993, so comparison of total Site 300 dose can only be made for 1993 and later. In addition, diffuse source doses were not reported separately from the total dose for the Livermore site for 1990 and 1991.

Doses from Unplanned Releases

There were no unplanned atmospheric releases of radionuclides at the Livermore site or Site 300 in 2002.

Table 4. List of facilities or sources whose emissions accounted for more than 90% of the SW-MEI doses for the Livermore site and Site 300 in 2002.

Facility (source category)	CAP88-PC Dose in mrem/y	CAP88-PC Percentage contribution to total dose
Livermore site		
Building 612 Yard (diffuse source)	0.011*	48%
Building 331 stacks (point source)	0.0081*	35%
Building 514 Evaporator (point source)	0.0012	5.2%
Building 612, R102 (point source)	0.0011	4.8%
Building 331 Outside (diffuse source)	0.00087*	3.8%
Site 300		
Building 851 Firing Table (point source)	0.018	85%
Soil resuspension (diffuse source)	0.0033	15%

^{*} When LLNL's NEWTRIT model (see Section VII, subsection on "Modeling dose from tritium") is used in CAP88-PC in place of CAP88-PC's default tritium model, the doses for the diffuse Building 612 yard and Building 331 Outside sources are reduced to 0.75 of the values shown, and that for the Building 331 stacks is reduced to 0.69 of the value shown. Doses for other sources in the table are practically unchanged, since they have minor contribution from tritium.

Population Doses

Population doses, or collective EDEs, for both LLNL sites were calculated out to a distance of 80 km in all directions from the site centers using CAP88-PC. As noted earlier, CAP88-PC evaluates the four principal exposure pathways: ingestion through food and water consumption, inhalation, air immersion, and irradiation by contaminated ground surface.

Population centers affected by LLNL emissions include the relatively nearby communities of Livermore and Tracy, the more distant metropolitan areas of Oakland, San Francisco, and San Jose, and the San Joaquin Valley communities of Modesto and Stockton. Within the 80 km outer distance specified by DOE, there are 6.9 million residents included for the Livermore site population dose determination, and 6.0 million for Site 300. Population data files (distribution of population with distance and direction) used for the present report were the same as in the previous two years; see Tables 7 and 8 in *LLNL NESHAPs* 2000 Annual Report (Gallegos et al. 2001).

The CAP88-PC result for potential population dose attributed to 2002 Livermore-site operations was 0.50 person-rem (0.0050 person-Sv); the corresponding collective EDE from Site 300 operations was 2.5 person-rem (0.025 person-Sv). These values are both quite small and within the normal range of variation seen from year to year. By way of comparison, the population dose in the United States from exposure to the average level of natural background radioactivity is 1.9×10^6 person-rem (1.9×10^4 person-Sv).

Table 5. Doses (in mrem) calculated for the Site-Wide Maximally Exposed Individual (SW-MEI) for the Livermore site and Site 300, 1990 to 2002.

Year	Total Dose	Point Source Dose	Diffuse Source Dose
Livermore site			
2002	0.023 a	0.010 a	0.013
2001	0.017 a	0.0057 ^a	0.011
2000	0.038 a	0.017 a	0.021
1999	0.12 a	0.094 a	0.028
1998	0.055 a	0.031 a	0.024
1997	0.097	0.078	0.019
1996	0.093	0.048	0.045
1995	0.041	0.019	0.022
1994	0.065	0.042	0.023
1993	0.066	0.040	0.026
1992	0.079	0.069	0.010
1991	0.234	— b	— b
1990	0.240	− b	— b
Site 300			
2002	0.021	0.018	0.0033
2001	0.054	0.050	0.0037
2000	0.019	0.015	0.0037
1999	0.035	0.034	0.0012
1998	0.024	0.019	0.005
1997	0.020	0.011	0.0088
1996	0.033	0.033	0.00045
1995	0.023	0.020	0.003
1994	0.081	0.049	0.032
1993	0.037	0.011	0.026
1992	0.021	0.021	<u>—</u> с
1991	0.044	0.044	— с
1990	0.057	0.057	— с

^a The dose includes HT emissions modeled as HTO as directed by EPA Region IX. EPA Region IX acknowledges that such modeling results in an overestimation of the dose. This methodology is used for purposes of compliance.

Compliance with 40 CFR 61 Subpart H (61.93)

Calculations of effective dose equivalents for Livermore-site and Site 300 facilities having the potential to release radioactive material to the atmosphere were found to be well below the 10 mrem (100 μ Sv) NESHAPs dose standard for dose to the most-exposed individual members of the public. Tritium accounted for more than 87% of the Livermore-site calculated dose, while at Site 300 practically the entire calculated dose was due to the isotopes ²³⁸U, ²³⁵U, and ²³⁴U, in depleted uranium.

b Diffuse source doses were not reported separately from the total dose for the Livermore site for 1990 and 1991.

^c No diffuse emissions were evaluated at Site 300 for years before 1993.

In 2002, there were seven buildings (Buildings 175, 177, 235, 251, 331, 332, and 491) at the Livermore site and one (Building 801, the Contained Firing Facility) at Site 300 that had radionuclide air effluent monitoring systems. These buildings are listed in Table 2, along with the number of samplers, the types of samplers, and the analytes of interest.

LLNL remains committed to monitoring stack effluent air from its Tritium Facility (Building 331), Plutonium Facility (Building 332), Contained Firing Facility (Building 801), and the seismically hardened area of its Heavy Element Facility (Building 251). In addition, other facilities are continuously monitored, as necessary, based on evaluations of potential emissions without control devices, as in the case of Building 235, or where classification or other issues prevent a usage-inventory-based evaluation.

SECTION V. Certification

I certify under penalty of law that this document and all attachments were prepared under my direction or supervision in accordance with a system designed to assure that qualified personnel properly gather and evaluate the information submitted. Based on my inquiry of the person or persons who manage the system, or those persons directly responsible for gathering the information, the information submitted is, to the best of my knowledge and belief, true, accurate, and complete. I am aware that there are significant penalties for submitting false information, including the possibility of fine and imprisonment for knowing violations.

Name:	Dennis K. Fisher
	Associate Director Safety and Environmental Protection
	Lawrence Livermore National Laboratory
	7000 East Avenue, L-668
	Livermore, CA 94550
Signature:	Date:
	Dennis K. Fisher
the informati immediately information i penalties for	er penalty of law that I have personally examined and am familiar with on submitted herein, and based on my inquiry of those individuals responsible for obtaining the information, I believe that the submitted is true, accurate, and complete. I am aware that there are significant submitting false information, including the possibility of fine and it. See 18 U.S.C. 1001.
Name:	Phillip Hill Acting Deputy Manager Safety and Environmental Programs U.S. Department of Energy 7000 East Avenue, L-293 Livermore, CA 94550
Signature:	Date:
	Phillip Hill

SECTION VI. Supplemental Information on NESHAPs Compliance and QA/QC Activities

Requirements Under New EPA Standard for Stack Sampling

In September 2002 EPA amended 40 CFR 61 Subpart H (NESHAPs) to require use of a new standard, ANSI N13.1-1999, for stack sampling of radiological effluent from certain newly constructed or modified facilities. This action replaced the existing standard ANSI N13.1-1969, and imposed some conditions on stack monitoring systems of existing facilities that are "grandfathered in" under the old standard. An assessment performed by TAMM Group in EPD identified 10 stack sampling systems (nine at the Livermore site and one at Site 300) that must satisfy the new standard, as listed in the following table.

Table 6. Livermore site and Site 300 stack sampling systems that must satisfy the maintenance and inspection requirements in the ANSI N13.1-1999 standard.

Building	Exhaust	Sampler ID	Operation	
251	FGBE-1000	PAM_46	Hardened Area Glove Boxes	
251	FGBE-2000	PAM_47	Hardened Area Glove Boxes	
695 ^(a)	FHE-1000, 2000, 3000	PAM_1	FHE, Waste Treatment Exhaust	
332	FGBE-1000	SP_3	Glove Box, Increment 1	
332	FGBE-2000	SP_4	Glove Box, Increment 1	
332	FGBE-3000	SP_8	Glove Box, Increment 1	
332	FGBE-4000	SP_9	Glove Box, Increment 1	
332	FGBE-7000, 8000	SP_10	Glove Box, Increment 3	
801	FEFH-1, FE-2	PAM_1	Test Chamber, Facility Exhaust	
235	FHE-2001, 2002	PAM_1	Hood and Glove Box Exhaust, Room 1130	

^a The stack for Building 695, LLNL's new Decontamination and Waste Treatment Facility, was not operational in 2002.

An implementation plan was prepared that addresses the inspection and calibration requirements of the new standard. The LLNL stack monitoring systems not cited in Table 6 are not required by NESHAPs regulations, but continue in operation as a best management practice. The new standard is described in a 1999 supplement to Health Physics Society Journal, entitled "Sampling and monitoring releases of airborne radioactive substances from the stacks and ducts of nuclear facilities" (report ANSI/HPS N13.1-1999).

Periodic Confirmatory Measurements

Results of NESHAPs periodic confirmatory measurements serve to support or confirm two objectives: (1) that those operations not continuously monitored do not, in fact, need to be continuously monitored, and (2) that radionuclide usage-inventory-based estimates of emissions and their corresponding doses are conservative. The particular sampling system chosen for study was randomly selected from a set of significant candidate sampling operations.

In 2002, periodic confirmatory sampling was conducted for a period of two weeks at Building 151, focused specifically on a dual stage HEPA filter system. This system ventilates a glovebox in which liquid samples of weapons grade Pu-238 are chemically purified. All measured concentrations were less than the minimum detectable concentrations for alpha and beta activity. Projecting these results to occur for an entire calendar year yields potential SW-MEI doses that would be seven orders of magnitude less than the EDE from all Livermore site operations for 2002. Because the dose calculated is estimated from a minimum detectable emission rather than an actual measured emission, it represents an upper-bound dose estimate, and is consistent with the dose based on the inventory approach and reported in Attachment 1.

Proposal to EPA for Use of Surveillance Air Monitoring in Demonstrating NESHAPs Compliance for LLNL's Numerous Minor Sources

In 2002 LLNL drafted a proposal to EPA Region IX, requesting permission to use surveillance air monitoring data in demonstrating NESHAPs compliance of radiological releases from the nearly 200 minor stack and diffuse sources at the Livermore site. This data would be used in place of inventory-based modeling, resulting in savings in time and money to both EPD and Laboratory Programs. This proposal and EPA's response in accepting it are reproduced in Attachment 3.

NESHAPs Quality Assurance (QA) Program

The LLNL NESHAPs quality assurance program is a multi-organizational effort that is described in the *Lawrence Livermore National Laboratory Quality Assurance Project Plan for National Emission Standards for Hazardous Air Pollutants (NESHAPs), 40 CFR 61, Subpart H (QAPP—Hall, L.C. and A.H. Biermann, UCRL-ID-13914, 2000).* The QAPP is structured in the manner prescribed for quality assurance programs that is outlined in Appendix B, Method 114 of 40 CFR 61. The QAPP describes the organization structure and functional responsibilities, objectives of the quality assurance program, administrative controls in place for handling sample collection systems, sample collection and effluent flow rate measurement systems, corrective actions, and reporting.

The major components of this multi-organizational effort are the LLNL facilities/programs that have continuous monitoring systems, the Radiological Measurements Laboratory (RML) and the Analytical Laboratory (AL), both in the Hazards control Department (HCD), and the Environmental Protection Department (EPD). In addition to the QAPP, NESHAPs Agreement of Roles and Responsibilities (NARRs) documents are in place between EPD and the facilities and/or programs and HCD; these NARRs formalize responsibilities and obligations of the organizations regarding many tasks for the air effluent sample network. Tasks that are addressed in the NARRs include air sampler design and installation, procedures and their implementation, sampling, sample analysis and tracking, maintenance and repair of sampling systems, guidance on regulatory requirements, documentation of the sampling network, reporting, and the archival of records.

EPD is responsible for an annual assessment and demonstration of LLNL's compliance with NESHAPs. The Department operates under a Quality Assurance Management Plan and associated procedures and guidance documentation. The Terrestrial and Atmospheric Monitoring and Modeling Group (TAMM) of EPD is responsible for environmental monitoring; air dispersion and dose assessment modeling; assessment (in cooperation with Laboratory Program personnel) of usage and potential release of radioactive materials to air in operations throughout the Laboratory; and reporting to EPA and DOE to demonstrate the Laboratory's compliance with NESHAPs. Detailed records are kept of all measurements, computer model runs and other calculations, and selected model runs are validated. The TAMM group is informed of proposed new operations, and modified operations where significant changes in radiological usage inventories occur, by several mechanisms. These include reviews of National Environmental Policy Act (NEPA) documentation, Integration Worksheets, Occupational Safety Plans (describing facility-specific safety procedures and plans), and knowledge derived from participation on EPD's Environmental Support Teams. All NESHAPs evaluations and calculations, along with supporting information, are archived for at least the period of time specified in 40 CFR 161 Subpart 1H.

Quality Control (QC) for 2002 Radiological Usage Inventory and Modeling

Of the four-dozen sources for which modeling runs were performed in the reduced accounting for 2002, approximately 15% were selected for validation, which entails confirmation of both the source emission data and dose modeling calculations. Two sources (one from each of the two LLNL sites) were selected because they represented the most significant contributions to 2002 potential dose to the public; five additional sources were selected as the most important radiological activities in Radioactive and Hazardous Waste Management (RHWM) Division (from a public dose standpoint); and one significant diffuse source was selected. Specifically, the sources chosen for quality control review were the following: the Tritium Facility's

two 30-m stacks; one explosives experiment conducted at Site 300's Firing Table 851; five sources reported by RHWM; and the Building 612 Yard waste tritium storage area.

More broadly, the quality and accuracy of our accounting and inventory processes were checked in several ways. In the accounting of new sources, more than 200 NEPA or related (primarily Integration Work Sheets and Occupational Safety Plans) documents were examined as they arose over the course of the year and reexamined collectively at year's end to identify all new 2002 projects having potential to release radioactive material to air. Additionally, all Radioactive Materials Management Areas new to 2002 were inventoried. The data characterizing the principal source at each site (principal in terms of producing the greatest potential dose to the public) were double-checked for accuracy. Finally, each radiological inventory form returned by the programs was scrutinized for consistency and evident errors as it was compiled and entered into the spreadsheet, Attachment 1. Based on these QC efforts, we believe that the data presented in Attachment 1 meets EPD's quality assurance objectives.

SECTION VII: Supplementary Information on Radiological Dose Assessment for 2002

Livermore-Site Principal Diffuse Sources

The dose evaluations for diffuse sources at the Livermore site in 2002 required several different modeling approaches. Building 331 Outside Yard and Building 612 Yard emissions estimates were based on facility personnel knowledge and environmental surveillance data. Building 514 Tank Farm emissions estimates were derived from radiological usage inventory data. The dose in each of these cases was calculated using CAP88-PC. Air surveillance monitoring data for plutonium from a monitor located at the SW-MEI was used to evaluate the dose from plutonium contamination in the Southeast Quadrant.

Building 331 Outside Yard

As the Tritium Facility (Building 331) conducts operations, tritium-contaminated equipment and material slated for disposal is removed from the building, packaged in a waste accumulation and storage area, removed from the building to an outside storage container, and sent to Radioactive and Hazardous Waste Management Division (RHWM) facilities. During 2002, outgassing from such waste released an estimated 1.0 Ci (3.7 \Box 10¹⁰ Bq) of tritium to the atmosphere outside Building 331. This amount was derived from process and facility knowledge and environmental surveillance measurements. This release was modeled in CAP88-PC as a 1 m² area source, leading to a calculated 2002 dose to the SW-MEI of 8.7 \Box 10⁻⁴ mrem (8.7 \Box 10⁻³ μ Sv); a dose 0.75 times this amount was calculated when the NEWTRIT model was implemented in CAP88-PC.

Building 514 Tank Farm

Another potential source of diffuse emissions of a variety of radionuclides was RHWM waste storage and treatment operations. Building 514 houses the RHWM "Tank Farm," consisting of six 7,170-liter tanks with ancillary equipment such as pumps, mixers, probes, and a bulking station. The tanks are used to store and treat liquid and solid radioactive and/or mixed wastes. Treatment is performed on a batch basis. Chemicals and waste are added to the tanks to achieve the desired treatment objectives. A 2002 radionuclide usage inventory was conducted for the facility to determine the diffuse source term (Attachment 1). CAP88-PC modeling gave a 2002 SW-MEI dose for the Tank Farm to be 4.1 \square 10⁻⁴ mrem (4.1 \square 10⁻³ μ Sv).

Building 612 Yard

The Building 612 Yard is a potential source of diffuse emissions of tritium. This area is dedicated to hazardous waste, radioactive waste, and mixed waste management activities. The yard consists of several areas where waste containers are stacked outdoors. Several of these containers outgas tritium. A surveillance air monitor designated B624 has been placed in the Building 612 Yard to provide continuous

measurements of tritium in air near this source. The median annual concentration of tritium in air for 2002 in this area was 49 pCi/m³ (1.9 Bq/m³). These data were used to calculate the total tritium emissions from the area, using a conservative approach that assumed the source to be 60 m south-southwest of the air sampler. With this assumption, a diffuse source emission of 2.3 Ci/y (7.4 \Box 10¹⁰ Bq/y) was required to produce the concentrations measured at the air sampler. This source term produced a CAP88-PC-calculated 2002 dose to the SW-MEI from the Building 612 Yard of 1.1 \Box 10⁻² mrem (1.1 \Box 10⁻¹ μ Sv); a dose 0.75 times this amount was calculated when the NEWTRIT model was implemented.

Southeast Quadrant

The Southeast Quadrant of the Livermore site has elevated levels of plutonium in the surface soil (from historic waste management operations) and air (from resuspension). A high volume air particulate sampler is located adjacent to the UNCLE Credit Union (the location of the SW-MEI) to monitor the plutonium levels in this area. Monitoring data from this air sampler were used as a direct measurement of potential dose via the air pathway. The median annual concentration of $^{239+240}$ Pu (the analytical technique used, namely alpha spectroscopy, does not distinguish between 239 Pu and 240 Pu) in air was 1.83×10^{-19} Ci/m³ (6.76×10^{-9} Bq/m³). Using the dose conversion factor of 3.08×10^{-19} mrem/ μ Ci (8.32×10^{-5} Sv/Bq) from Federal Guidance Report No. 11, EPA-520/1-88-020, U.S. Environmental Protection Agency (1988) for 239 Pu and 240 Pu, and the standard man breathing rates of 8400 m^3 /y, the dose was determined to be 4.7×10^{-4} mrem ($4.7 \times 10^{-3} \mu$ Sv) for 2002.

Site 300 Principal Diffuse Sources

Diffuse sources at Site 300 involve primarily depleted uranium, and to a considerably lesser extent, tritium. During remediation efforts at Site 300, LLNL completed a contaminant screening to identify potential routes of migration from soil to air and other environmental media of these radionuclides and other contaminants (Final Site Wide Remedial Investigation Report; Webster-Scholten, Ed., 1994, Lawrence Livermore National Laboratory, Livermore, CA, UCRL-AR-108131). Uranium-238 and tritium were identified as contaminants of potential concern.

Tritium Evaporation and Migration at Site 300

Tritium gas and solids containing tritium (Li³H) were components of explosives assemblies tested on the firing tables during experiments in years past. Most of the gaseous tritium escaped to the atmosphere during the tests, but some of the solid Li³H remained as residue in the firing table gravel. Rainwater and dust-control rinse water percolated through the gravel, causing the tritium to migrate into the subsurface soil and, in some cases, eventually to the ground water. Tritium contaminated gravel was removed from the firing tables in 1988 and disposed in the Pit 7 landfill. Tritium in landfills, firing table soils, and ground water are potential sources of diffuse emissions of tritium to the atmosphere at Site 300. LLNL

personnel maintain an air tritium sampler at a perimeter location at Site 300, and doses from diffuse tritium sources may be estimated based on the monitoring data for that sampling location. For the calendar year 2002, all measurements in ambient air at the Site 300 perimeter location were consistent with natural background measurements.

Resuspension of Depleted Uranium in Soil at Site 300

Like tritium, depleted uranium has been used as a component of explosives test assemblies. It remains as a residue in surface soils, especially near the firing tables. Because surface soil is subject to resuspension by the action of wind, rain, and other environmental disturbances, the collective effects of surface soil uranium residuals on off-site doses were evaluated.

A model was developed to distinguish between the contribution to measured uranium activities arising from naturally occurring uranium (NU) and that from depleted uranium (DU) contributed by LLNL operations. (A derivation of the model was presented in *LLNL NESHAPs* 1995 *Annual Report*, Gallegos et al., 1996.) We base our dose estimate for resuspended depleted uranium (DU) on the measured environmental surveillance monitoring total concentration in air of uranium-238, subtracting out the part contributed by NU, from the following equation:

$$\Box = \frac{0.00726 \,\Box\, 0.99274 \, \frac{M(CU \,\Box\, 235)}{M(CU \,\Box\, 238)}}{0.00526 \, \frac{M(CU \,\Box\, 235)}{M(CU \,\Box\, 238)} + 0.00526}$$

where μ is the fraction (by weight) of uranium contributed by operations, CU is composite uranium (both DU and NU), M(CU-235) the mass of U-235 in the composite (measured) uranium, and M(CU-238) the mass of U-238 in the composite (measured) uranium.

For 2002, all eight air-particulate monitors at Site 300 were used to determine the annual-average concentrations of isotopes U-238 and U-235. These site-average values gave an estimate of 3.3×10^{-3} fmrem ($3.3 \times 10^{-2} \mu Sv$) for the SW-MEI dose resulting from resuspension of DU in soil for 2002.

Modeling Dose from Tritium

To evaluate dose from tritium releases to air, we use the EPA-sanctioned CAP88-PC code. Its tritium model calculates dose from inhalation, skin absorption, and ingestion of tritium only in its tritiated water vapor form (HTO). Doses from HT or organically bound tritium (OBT) are not calculated. CAP88-PC's tritium model is based on the specific activity model, which assumes that the tritium-to-hydrogen ratio in body water is the same as in air moisture. Because the specific activity model

is linked in CAP88-PC with relatively high dose coefficients for HTO, the model's dose predictions generally err on the high side.

Inhalation doses from unit concentration of HT in air are a factor of 15,000 times lower than those from inhalation and skin absorption of unit concentration of HTO in air (International Commission on Radiological Protection (ICRP), 1995, *Age dependent doses to members of the public from intake of radionuclides, Part 4, Inhalation Dose Coefficients*. Oxford: Pergamon Press; ICRP Publication 71; Ann. ICRP 25[3&4]). Thus, doses from inhaled HT can safely be ignored unless the air concentration is extremely high. A release of HT cannot be ignored, however, because HT that reaches the ground is rapidly and efficiently converted to HTO by microorganisms in soil (McFarlane, Rogers, and Bradley, Environmental Science and Technology 12: 590-593,1978; Brown, Ogram, and Spencer, Health Physics 58:171-181, 1990) and to a lesser extent in vegetation (Sweet and Murphy, Environmental Science and Technology, 18:358-361, 1984).

Organically bound tritium (OBT) is formed by plants during photosynthesis and is incorporated by animals when ingested. Animals also metabolize some OBT from ingested or inhaled HTO. The ICRP dose coefficient for OBT is about 2.3 times higher than that of HTO, because the biological half-life of OBT in the body is longer than that of HTO, which is eliminated at the same rate as body water. Although doses predicted by CAP88-PC are generally high enough to account for dose from ingested OBT, nevertheless, a model that explicitly calculates dose from OBT is preferable.

A simple tritium model, NEWTRIT, has been developed that calculates ingestion dose from both HTO and OBT and accounts for conversion of HT to HTO in the environment following releases of HT (Peterson, S-R. and P.A. Davis, Health Physics 82(2):213-225, 2002). For calculating doses in this report, LLNL has used the NEWTRIT model in CAP88-PC, in addition to the default CAP88-PC code, to estimate doses from significant sources of tritium emissions; see, e.g., Table 4. A brief discussion of the NEWTRIT model was presented in Attachment 2 of the 2000 NESHAPs annual report (*LLNL NESHAPs 2000 Annual Report*, Gallegos et al. June 2001).

In October 2001, LLNL sent a letter to EPA Region IX requesting consideration of NEWTRIT as an alternative methodology for calculating doses from atmospheric releases of tritiated water vapor (HTO) and tritiated gas (HT), for use in demonstrating compliance with radionuclide NESHAPs (40 CFR 61 Subpart H). In late 2002, the EPA had NEWTRIT coded into GENII-NESHAPs, a version of GENII (Napier et al. 1988) that the EPA plans to approve as a regulatory model for evaluating radionuclide NESHAPs compliance. At this writing, GENII-NESHAPs is undergoing peer review.

Comparison of 2002 Modeling Results with Tritium Air Surveillance Monitoring Data

A comparison was made between CAP88-PC-predicted concentrations of tritium in air and ambient air monitoring data for eleven tritiated water vapor samplers on the Livermore site (designated VIS, SALV, POOL, CAFE, MESQ, MET, COW, B331, B514, B624, and B292) and one off-site sampler (ZON7) that have been used for comparison since 1997. In addition, a new air tritium monitor (DWTF) has been added to the comparison. Monitor locations are shown in Figure 7.

Only concentrations from the three most significant sources of tritium releases to air at the Livermore site were included in the model-data comparison. The largest point source is the Tritium Facility (Building 331), where tritium is emitted from two 30-mhigh, continuously monitored stacks. Based on stack monitoring, a total of 32.9 Ci $(1.22 \,\square\, 10^{12}\,\text{Bg})$ of HTO was emitted from Building 331 stacks in 2002. (The 3.47 Ci [1.28] $\Box 10^{11}$ Bq] of HT emitted from the Tritium Facility stacks is not included in the comparison because the tritium air surveillance monitors register only HTO.) Generally one would expect the Tritium Facility stacks to make the largest contribution to concentrations of tritium at distant monitors (e.g., ZON7), because the emissions are cast high into the air and carried with the wind. Diffuse-source emissions are lower to the ground, primarily affecting those monitors in close proximity. The other two principal sources in our modeling/measurement comparison are of this type: open-air diffuse emission areas associated with the Building 612 Yard and the Tritium Facility (Building 331) outside yard waste accumulation and storage areas. Emissions from the Building 612 Yard source were estimated to be 2.3 Ci (8.5 ☐ 10¹⁰ Bq), based on calibrating CAP88PC-predictions of tritium concentrations at the tritium monitor B624 closest to it. (Thus the B624 data do not provide a test of the modeling.) Emissions from the B331 outside yard source were estimated to be 1.0 Ci (3.7 \square 10¹⁰ Bq) in 2002, based on facility knowledge and environmental monitoring data (primarily the B331 monitor near this yard). While these two diffuse sources contribute significantly to tritium concentrations in all of the monitors, all other potential sources of tritiated water vapor release, such as the radioactive and hazardous waste management operations in Building 514 and the Building 292 diffuse source, were too minor to influence the overall model-data comparison.

Annual average concentrations of HTO in air (pCi/m^3) at the locations of the thirteen monitors were modeled for the three sources individually and collectively, and compared to the measured annual mean concentrations . The results, displayed in Table 7, show that by taking into account the leading sources releasing tritiated water vapor to air, fairly good agreement is obtained between model runs and data for all of the air tritium monitors.

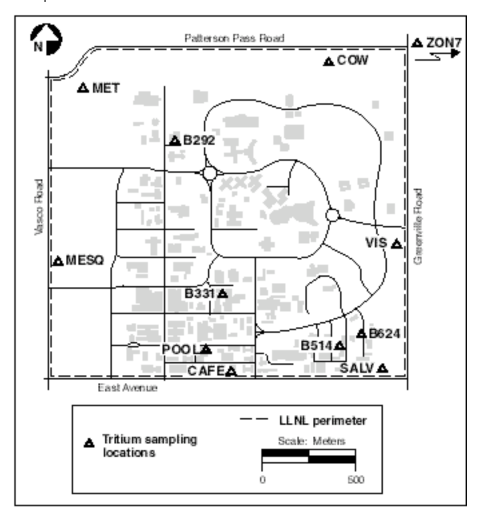


Figure 7. Tritiated water vapor surveillance sampling locations, Livermore site.

With the exception of the monitor B292 (which is probably under-predicted due to our neglect of a small contribution from a diffuse source nearby that only impacts that monitor), all predictions are equal to or greater than what was measured at the monitors. This means that in 2002, as in the past, CAP88-PC is over-estimating HTO in air from LLNL releases of HTO, particularly because CAP88-PC cannot account for the small amount of HTO resulting from conversion of HT. This consistent bias towards over-estimation since 1997 is probably caused by the relative importance of the diffuse sources for these years (Peterson, S-R. Testing CAP88-PC's Predicted Air Concentrations Against Historical Air Tritium Monitoring Data, 1986 – 2001, at Lawrence Livermore National Laboratory. In draft, 2003.). A comparison of AIRDOS-EPA predictions of air concentrations for various radionuclides (²³⁴U, ²³⁸U, ⁸⁵Kr, and ³H) with measurements at six different sites concluded that the 90% confidence interval for the accuracy of the CAP88-PC dispersion model ranged from a factor of 0.3 to 4.4, based on 51 samples ("Comparison of AIRDOS-EPA predictions of ground-level airborne radionuclide concentrations to measured values," Jack

Faucett Associates, Bethesda, MD. 20814; JACKFAU-341/12-87; 1987). Similarly, the study (Peterson op. cit.) that compared CAP88-PC predictions with air tritium concentrations at 13 perimeter and off-site locations showed that ninety-six percent of all predictions fell within a factor of three of the observations, and slightly more than half of the predicted air concentrations were greater than the observed air concentrations.

Table 7. Comparison of measured and modeled annual mean concentrations of tritiated water vapor (HTO) in air at selected Livermore site locations, 2002.

Air monitor (name)		Modeled* average concentration (pCi/m ³)	Ratio of modeled- to-measured concentrations	Modeled of tritium in the indicate		buted by
				B331	B612	B331
				Stacks	Yard	Outside
B624	56.4	58	1.0	1.4	56	0.12
B331	10.0	14	1.4	0.051	1.4	13
POOL	3.22	3.5	1.1	1.2	1.2	1.1
B514	3.15	8.4	2.7	0.56	7.7	0.11
B292	1.75	0.77	0.46	0.23	0.32	0.22
VIS	1.72	2.6	1.5	1.2	1.3	0.14
CAFE	1.67	2.2	1.3	0.68	1.2	0.35
DWTF	1.45	1.5	1.0	1.2	0.24	0.10
COW**	1.22	1.4	1.1	1.0	0.24	0.12
SALV**	0.929	1.6	1.7	0.40	1.1	0.061
MESQ**	0.755	0.97	1.3	0.20	0.35	0.42
ZON7**	0.663	0.67	1.0	0.50	0.14	0.032
MET**	0.458	0.49	1.1	0.15	0.19	0.15
(CRED)***		3.5		1.3	2.0	0.16

^{*}This result takes into account the three most significant tritium sources; it is the annual-average concentration comprising the sum of the three contributions shown in the far right columns.

^{**}At these locations, more than 25% of the samples were below detection limits. The annual mean includes negative concentrations for all except COW. MET has the lowest percentage of detections (17%).

^{***}The CRED location does not have a tritium surveillance air monitor, but it marks the location of the SW-MEI.

SECTION VIII. Supplemental Information on Other Compliance

Status of Compliance with Other Regulations
Status of compliance with 40 CFR 61 Subpart Q - National Emission
Standards for Radon Emissions from Department of Energy Facilities
LLNL does not have storage and disposal facilities for radium containing materials that would be a significant source of radon.

Status of compliance with 40 CFR 61 Subpart T - National Emission Standards for Radon Emissions from the Disposal of Uranium Mill Tailings LLNL does not have or store any uranium mill tailings.

Information on Radon-220 and Radon-222 Emissions

Radon emissions occur naturally by emanation from the earth. Radon-222 emissions that were reported in past NESHAPs annual reports from research experiments at the Livermore site did not occur in 2002.

ATTACHMENT 1. LLNL NESHAPs 2002 Annual Report Spreadsheet

Guidance for Interpreting the Data Spreadsheet

A generalized description of each facility and its operations is provided on the spreadsheet. In addition, the following information is shown for each listed emission point or stack:

- Building and room number(s)
- Specific stack identification code(s)
- Generalized description of operations in the room(s) or area(s)
- Radionuclides utilized in the operation
- Annual radionuclide usage inventory with potential for release (by isotope, in curies)
- Physical state factors (by isotope)
- Stack parameters
- Emission control devices and emission control device abatement factors
- Estimated or measured annual emissions (by isotope)
- Distance and direction to the site-wide maximally exposed individual (SW-MEI)
- Calculated EDE to the SW-MEI
- Distance and direction to the maximally exposed individual for that specific source (MEI)
- Calculated EDE to the MEI (source term not adjusted for emission controls)
- Source category

Radionuclides

The radionuclides shown in the spreadsheet are those from specific emission points where air emissions were possible. If radionuclides were present, but encapsulated or sealed for the entire year, radionuclides, annual usage inventories, and emissions are not listed.

Radionuclide Usage Inventories with Potential for Release

The annual radionuclide usage inventories for point source locations are based on data from facility experimenters and managers. For Buildings 251 (hardened area) and 332, classification issues regarding transuranic radionuclide usage inventories make use of the usage inventory/modeling approach impractical. However, all such affected emission points in these buildings are continuously monitored, and emissions are therefore directly determined.

Physical State Factors

The physical state factors listed are EPA potential release fractions from 40 CFR 61, Appendix D, whereby emissions are estimated from radionuclide usage inventories depending on their physical states for use in dispersion/dose assessment modeling. A physical state factor of $1.0 \,\square\, 10^{-6}$ is used for solids, $1.0 \,\square\, 10^{-3}$ is used for liquids and powders, and 1.0 is used for unconfined gases and substances heated above 100° C. Regarding the latter, U.S. EPA has granted LLNL approved alternative emissions factors for elemental uranium, uranium/niobium alloy, and elemental plutonium. (See Table 3 in Section III.) These factors are allowed provided that the material is not intentionally dispersed to the environment and that the processes do not alter the chemical form of the material.

Stack Parameters

Engineering surveys conducted from 1990 through 1992 form the basis for the stack physical parameters shown, which were checked and validated by facility experimenters and managers for 1994 and 1995. Stack physical parameters for sources evaluated in 2002 were updated, as necessary, by experimenters and managers for those facilities.

Emission Control Devices

High Efficiency Particulate Air (HEPA) filters are used in many LLNL facilities to control particulate emissions. For some discharge points, scrubbers and electrostatic precipitators aid the control of emissions. The operational performance of all HEPA filtration systems is routinely tested. The required efficiency of a single stage HEPA filter is 99.97%. Double staged filter systems are in place on some discharge points. Triple stage HEPA filters are used on glove box ventilation systems in the Building 332 Plutonium Facility and in the hardened portion of Building 251.

Control Device Abatement Factors

Similar to physical state factors, control device abatement factors, from Table 1 in 40 CFR 61, Appendix D, are those associated with the listed emission control devices, and are used to better estimate actual emissions for use in dispersion and dose models. By regulation, each HEPA filter stage is given a 0.01 factor (even though the required test efficiency that all LLNL HEPA filters must maintain would yield a factor of 0.0003).

Estimated Annual Emissions

For unmonitored and non-continuously monitored sources, estimated annual emissions for each radionuclide are based on the product of (1) usage inventory data, (2) time factors (discussed in "Emission Source Terms" in Section III, (3) EPA potential release fractions (physical state factors), and (4) applicable emission control device abatement factors.

Actual emission measurements are the basis for reported emissions from continuously monitored facilities. LLNL facilities that had continuous monitoring systems in 2002 were Buildings 175, 177, 235, 251, 331, 332, and 491 at the Livermore site, and Building 801 at Site 300, as noted earlier. See the discussion below under "0.1 mrem/y Monitoring Requirement" regarding the use of emissions measurements for monitored sources.

10 mrem/y Site-Wide Dose Requirement

For LLNL to comply with the NESHAPs regulations, the LLNL site-wide maximally exposed individual (SW-MEI; defined as the hypothetical member of the public at a single residence, school, business, or office who receives the greatest LLNL-induced EDE from the combination of all radionuclide source emissions) cannot receive an EDE greater than 10 mrem/y (100 μ Sv/y). (See Section III for a discussion of the SW-MEI.)

In the spreadsheet, the distance and direction to the respective SW-MEI are shown for each facility at each site. Doses to the site specific SW-MEIs were evaluated for each source and then totaled for site specific evaluations against the 10 mrem/y dose standard (see Section IV).

0.1 mrem/y Monitoring Requirement

To assess compliance with the requirement for continuous monitoring (potential dose greater than $0.1 \, \mathrm{mrem/y} \, [1.0 \, \mu \mathrm{Sv/y}]$ to the maximally-exposed public individual or MEI, discussed earlier in Section III), emissions must be individually evaluated from each point source. The location of the MEI is generally different for each emission point. The maximum dose at a location of unrestricted public access typically occurs at a point on the site perimeter. Therefore, it is often referred to as the maximum "fence line" dose, although the off-site maximum dose could occur some distance beyond the perimeter. (This could happen, e.g., when the perimeter is close to a stack; however, for nearly all emission points at the Livermore site and Site 300, calculations show that ground level concentrations of radionuclides generally decline continuously beyond LLNL boundaries.) As stipulated by the regulations, modeling for assessment of continuous monitoring requirements assumed unabated emissions (i.e., no credit was taken for emission abatement devices, such as filters), but physical state factors and time factors were applied.

The unabated EDE cannot be calculated for HEPA-filtered facilities monitored for radioactive particles. Because the monitoring equipment is placed after HEPA filtration, there is no way to obtain an estimate for what the emissions might have been had there been no filtration. It is not reasonable to apply factors for the effects of the HEPA filters on the emission rate because most of what is measured on the HEPA filters is the result of the radioactive decay of radon, which is capable of penetrating the filter. The spreadsheet gives, for each inventoried point source, the

dose to the MEI and the distance and direction to the LLNL fence line where the MEI is located. However, for HEPA-filtered monitored sources, no value is shown.

Source Categories

LLNL radionuclide air emission sources have been classified into seven source categories, indicated by the number in the next to last column of the spreadsheet: (1) Unmonitored or non-continuously monitored Livermore-site facilities that have had a radionuclide usage inventory update for 2002; (2) Unmonitored or non-continuously monitored Livermore site facilities with a previous radionuclide usage inventory update (this category is not used in years with complete usage inventory updates, such as 2000); (3) Continuously monitored Livermore site facilities; (4) Site 300 explosives experiments; (5) Diffuse sources where emissions and subsequent doses were estimated using inventory processes; (6) Diffuse sources where emission and dose estimates were supported by environmental surveillance measurements; and (7) Sources whose emissions estimates and subsequent doses were estimated by confirmatory air sampling rather than continuous sampling.

Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated			se Requirement		y Monitoring		Source
					with Potential for	State	Height (m)	Diameter	Velocity	Device(s)	Abatement	Annual Emissions	Distance to		EDE	Distance	Direction	Unabated	Category
IVEDMODE	E SITE POINT SOURC	rec			Release (Ci)	Factor		(m)	(m/s)		Factor	(Ci)	SWMEI (m)	to SWMEI	(mrem)	to MEI (m)	to MEI	EDE (mrem)	
LIVERMORE	E SITE POINT SOURC	LO																	
Building 13	1 complex is a large	office/laboratory facility hous	ing both Mechanical and Electrical Engir	neering Divisions.															
131	1221	FFE-02	Storage and cleaning of	U-238	6.1E-06	1.0E-06	12.2	0.15	7.8	HEPA	0.01	6.1E-14 7.9E-16	1326	E	3.1E-12	567	WNW	1.4E-09	2
			assemblies	U-235 U-234	7.9E-08 5.7E-07	1.0E-06 1.0E-06						7.9E-16 5.7E-15							
				0-234	3.7 L-07	1.0L-00						3.7 L-13							
131	1248	Room Air	Storage and display of	U-238	1.5E-06	1.0E-06	NA	NA	NA	None	1	1.5E-12	1326	E	8.6E-11	524	W	1.4E-09	2
			post-test materials	U-235	2.0E-08	1.0E-06						2.0E-14							
				U-234	1.4E-07	1.0E-06						1.4E-13							
131	12404	Room Air	Character and display of	U-238	7.7E-07	1.0E-06	NA	NA NA	NA	Nama	1	7.7E-13	1326	-	4.3E-11	524	W	6.9E-10	2
131	1248A	ROOM All	Storage and display of post-test materials	U-235	9.9E-09	1.0E-06	INA	INA	INA	None	1	9.9E-15	1326	E	4.3E-11	324	VV	6.9E-10	
			post test materials	U-234	7.2E-08	1.0E-06						7.2E-14							
			of activities, including the Directorate				66												
laboratories	s in the Analytical &	Nuclear Chemistry Division and	d Chemistry and Chemical Engineering D	ivision; and NAI Dire	ectorate Forensic Scie	nces Center o	offices and laborat	tories.											
132N	2671	FHE-6000/7000	Mass spectrometry analysis	Pu-238	6.1E-07	1.0E-06	38.1	2.13	11.3	Double HEPA	0.0001	6.1E-17	1504	E	2.8E-15	1918	NE	3.8E-11	2
				Pu-239	4.5E-08	1.0E-06			1			4.5E-18	130.			1 .3.5			T
				Pu-240	1.1E-08	1.0E-06						1.1E-18							
				Pu-241	1.1E-07	1.0E-06						1.1E-17							
				Pu-242 Am-241	1.4E-12 1.1E-09	1.0E-06 1.0E-06						1.4E-22 1.1E-19							
				U-234	6.1E-12	1.0E-06						6.1E-19							
				0 23 1	0.12.12	1.02.00						0.12.22							
132N	2675	FHE-6000/7000	Preparation of aqueous	U-234	2.6E-18	1.0E-03	38.1	2.13	8.6	None	1	2.6E-21	1504	E	6.7E-16	481	SW	1.0E-15	2
			solutions for analysis	U-235	3.3E-16	1.0E-03						3.3E-19							
				U-238	4.6E-14	1.0E-03						4.6E-17							
		FHE-6000/7000	Analysis of aqueous solutions	U-234 U-235	1.3E-14 1.7E-12	1.0E+00 1.0E+00	38.1	2.13	8.6	HEPA	0.01	1.3E-16 1.7E-14	1504	E	3.3E-11	481	SW	5.2E-09	2
				U-238	2.3E-10	1.0E+00 1.0E+00						2.3E-12							
				0 230	2.32 10	1.02+00						2.32.12							
132N	2679	FHE-6000/7000	Preparation of aqueous	U-234	3.7E-17	1.0E+00	38.1	2.13	8.6	HEPA	0.01	3.7E-19	1504	E	1.3E-12	481	SW	2.0E-10	2
			solutions for analysis	U-235	1.5E-14	1.0E+00						1.5E-16							
				U-238	7.4E-12	1.0E+00						7.4E-14							
				Th-232	5.5E-13	1.0E+00						5.5E-15							-
132N	2685	FHE-6000/7000	Transfer and solvent extraction	Cs-137	8.8E-09	1.0E-03	38.1	2.13	8.6	None	1	9.0E-12	1504	F	3.8E-11	481	SW	6.2E-11	2
	2000	1112 0000/1000	of waste samples	Co-60	4.4E-10	1.0E-03	3311	20	0.0	110110		4.5E-13		_	0.02		0	0.22	
			for PCB analysis	Sr-90	4.8E-09	1.0E-03						4.9E-12							
				Th-228	3.4E-13	1.0E-03						3.5E-16							
				Th-230	1.0E-12 7.2E-14	1.0E-03						1.0E-15 7.4E-17							
				Th-232 Pu-238	7.2E-14 1.0E-11	1.0E-03 1.0E-03						7.4E-17 1.1E-14							
				Pu-239	4.4E-10	1.0E-03						4.5E-13							
				Pu-240	2.7E-10	1.0E-03						2.8E-13							
				Pu-241	2.4E-10	1.0E-03						2.5E-13							
				Am-241	2.4E-11 6.8E-12	1.0E-03 1.0E-03	-					2.5E-14 7.0E-15	-			-			-
-				U-234 U-235	6.8E-12 3.9E-13	1.0E-03 1.0E-03	 					7.0E-15 4.0E-16	 						+
				U-238	1.2E-12	1.0E-03						1.2E-15							+
					1														
132N	2689	FHE-6000/7000	Differential Scanning Calorimetry	U-238	5.1E-07	1.0E-06	38.1	2.13	11.2	None	1	5.1E-13	1504	E	8.3E-12	481	SW	1.3E-11	1
			of chemical mixtures	U-235	6.5E-09	1.0E-06	-					6.5E-15	-						
				U-234	4.7E-08	1.0E-06	+		-		+	4.7E-14	+						+
132N	2694	FHE-6000/7000	Transfer and solvent extraction	Cs-137	5.7E-09	1.0E-03	38.1	2.13	8.6	None	1	5.7E-12	1504	F	2.5E-11	481	SW	3.9E-11	2
. 32.14	2001	0000, 7000	of waste samples	Co-60	2.7E-10	1.0E-03	33.1	25	0.0	110110	· ·	2.7E-13	1304		2.02 11	101	311	0.02 11	 _
			for volatiles analysis	Sr-90	3.0E-09	1.0E-03						3.0E-12							
				Th-228	2.2E-13	1.0E-03						2.2E-16							
				Th-230	6.5E-13	1.0E-03						6.5E-16	-						-
-				Th-232 Pu-238	4.4E-14 6.5E-12	1.0E-03 1.0E-03	+		 		-	4.4E-17 6.5E-15	+						
				Pu-238 Pu-239	6.5E-12 2.9E-10	1.0E-03 1.0E-03	 					6.5E-15 2.9E-13	 			1			
				Pu-240	1.8E-10	1.0E-03						1.8E-13							
				Pu-241	1.5E-10	1.0E-03						1.5E-13							
				Am-241	1.5E-11	1.0E-03						1.5E-14							
				U-234	4.3E-12	1.0E-03	-					4.3E-15	-						
																			1
				U-235 U-238	2.5E-13 7.8E-13	1.0E-03 1.0E-03						2.5E-16 7.8E-16							

Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated	10 mrem/y S	Site-Wide Do	se Requirement	0.1 mrem	/y Monitoring	Requirement	Source
					with Potential for	State	Height (m)	Diameter	Velocity	Device(s)	Abatement	Annual Emissions	Distance to		EDE	Distance	Direction	Unabated	Categor
					Release (Ci)	Factor		(m)	(m/s)		Factor	(Ci)	SWMEI (m)	to SWMEI	(mrem)	to MEI (m)	to MEI	EDE (mrem)	
132N	2870	FHE-6000/7000	Preparation of urania and	U-234	3.5E-06	1.0E-03	38.1	2.13	11.2	None	1	3.5E-09	1504	E	1.5E-07	481	SW	2.4E-07	1
			thoria aerogels	U-235	1.5E-07	1.0E-03						1.5E-10							
				U-238	3.3E-06	1.0E-03						3.3E-09							
				Th-232	1.1E-06	1.0E-03						1.1E-09							
132S	2788	FHE-6000/7000	Transfer of uranium	U-238	5.7E-10	1.0E-03	4.6	1.22	8.9	None	1	5.7E-13	1504	E	2.1E-11	481	SW	7.4E-11	2
1323	2700	111E 0000/1000	Transier of dramam	U-235	7.3E-12	1.0E-03	7.0	1.22	0.5	None	'	7.3E-15	1304		2.72 11	701	344	7.4211	
				U-234	5.3E-11	1.0E-03						5.3E-14							
			ies nuclear and isotope sciences to a wi								characterization and	l analysis.							
Building 15	also contains the Cl	hemistry and Materials Scien	ces Environmental Services laboratory v	where samples of wa	aste streams and envi	ronmental me	edia (air, water, s	oil etc.) are analy	zed for their rad	lionuclide content.									
151	1033	FHE-2	Evaporation and transfer	Cm-248	3.5E-07	1.0E-03	12.8	0.41	7.8	None	1	3.5E-10	1308	E	4.4E-07	768	SW	1.1E-06	2
131	1033	THE Z	of solutions	Cm-246	8.3E-07	1.0E-03	12.0	0.41	7.0	None	'	8.3E-10	1300		7.72 07	700	3**	1.12.00	
			or colucione	U-233	1.9E-09	1.0E-03						1.9E-12							
				Np-237	3.5E-08	1.0E-03						3.5E-11							
				Pu-244	1.8E-11	1.0E-03						1.8E-14							
														_					
151	1034B	FGBE-5/6	Sample preparation	Am-241	2.7E-04 3.0E-03	1.0E-03	7.0	0.13	3.7	Double HEPA	0.0001	2.7E-11	1308	E	1.5E-07	540	W	1.3E-02	2
				Pu-238 Pu-239	4.7E-03	1.0E-03 1.0E-03						3.0E-10 4.7E-10				584	NWN	1.3E-02	
				Pu-239 Pu-240	1.0E-03	1.0E-03						1.0E-10							
				Pu-241	1.6E-02	1.0E-03						1.6E-09							
				Pu-242	4.7E-09	1.0E-03						4.7E-16							
				U-234	3.3E-08	1.0E-03						3.3E-15							
				Am-243	1.4E-04	1.0E-03						1.4E-11							
151	1034B	FGBE-7/8	Radiochemical analysis of	Cf-249	5.50E-05	1.0E-03	7.0	0.13	3.7	HEPA	0.01	5.5E-10	1308	E	6.3E-07	540	W	7.3E-04	1
			heavy element samples	Pu-238	3.00E-04	1.0E-03						3.0E-09							
				Am-243	1.00E-05	1.0E-03						1.0E-10							
151	1039	FHE-43	Transfer of solutions	Cs-137	4.6E-10	1.0E-03	12.8	0.46	11.3	None	1	4.6E-13	1308	E	3.9E-11	768	SW	7.6E-11	2
131	1033	THE 45	Transfer of solutions	Sr-90	3.0E-10	1.0E-03	12.0	0.40	11.5	TAOTIC		3.0E-13	1300	_	3.3L 11	700	3**	7.02 11	
				Gross alpha	3.2E-10	1.0E-03						3.2E-13							
151	1123	FHE-41	Evaporation and transfer	Pu-239	2.5E-14	1.0E-03	12.8	0.30	6.6	None	1	2.5E-17	1308	E	3.5E-15	768	SW	1.0E-14	2
			of solutions	U-238	2.6E-15	1.0E-03						2.6E-18				584	WNW	1.0E-14	
151	1131A	Room Air	Preparation of aqueous samples	H-3	4.5E-07	1.0E-03	NA	NA	NA	None	1	4.5E-10	1308	E	4.0E-13	540	W	5.8E-12	1
			for stable isotope analysis	Cs-137	4.5E-10	1.0E-03						4.5E-13							
151	1241	FHE-68	Radiochemical separations of	U-234	1.0E-07	1.0E-03	13.1	0.30	6.6	None	1	1.0E-10	1308	E	6.0E-09	584	WNW	2.0E-08	1
			uranium isotopes	U-235	1.0E-08	1.0E-03						1.0E-11							
151	1303	FHE-2000	Sample preparation	U-238	8.4E-13	1.0E+00	11.9	0.48	15.4	None	1	8.4E-13	1308	E	4.4E-08	1125	NNE	8.2E-08	2
			and analysis (ICP-MS)	U-235	3.9E-14	1.0E+00						3.9E-14							
				U-234	8.4E-13	1.0E+00						8.4E-13							
				U-233 Pu-239	1.9E-10 3.1E-10	1.0E+00 1.0E+00						1.9E-10 3.1E-10							
				U-238	8.4E-14	1.0E+00						8.4E-17							+
				U-235	3.9E-15	1.0E-03						3.9E-18							
				U-234	8.4E-14	1.0E-03						8.4E-17							
				U-233	1.9E-11	1.0E-03						1.9E-14							
				Pu-239	3.1E-11	1.0E-03						3.1E-14							
151	1304 8 13044	FUE 2000	Description of according to 105 (12)	0- 107	F 05 07	1.05.00	110	0.40	15.4	NI	-	F 05 07	1200	_	2.15.07	1400	N.E	2.25.07	
151	1304 & 1304A	FHE-2000	Preparation of samples for ICP/MS;	Cs-137 Pu-239	5.0E-07 6.1E-09	1.0E+00 1.0E-03	11.9	0.48	15.4	None	1	5.0E-07 6.1E-12	1308	E	2.1E-07	1469	NE	3.3E-07	1
			analysis by ICP/MS	U-239 U-233	1.0E-08	1.0E-03 1.0E-03	 				1	1.0E-12				1			+
				U-234	1.0E-06	1.0E-03	<u> </u>					1.0E-11							+
				U-235	1.0E-11	1.0E-03		1				1.0E-14							
				U-238	1.0E-13	1.0E-03						1.0E-16							
151	1306A	FHE-1000	Acid cleaning of glassware	Cs-137	2.5E-08	1.0E-03	13.1	0.46	4.6	None	1	2.5E-11	1308	E	3.8E-10	584	WNW	1.2E-09	1
				Pu-239	2.5E-09	1.0E-03	1	1				2.5E-12							
				U-233	2.5E-11	1.0E-03		-				2.5E-14				1			
				U-234	2.5E-12	1.0E-03	-	+				2.5E-15	-			1			
				U-235 U-238	2.5E-12 2.5E-13	1.0E-03 1.0E-03						2.5E-15 2.5E-16							
				U-238	2.3E-13	1.UE-U3	 					2.3E-10				1			+
	1				1		1	1								1	1		+ -
151	1318	FHF-26	Sample preparation	Pu-239	1 0F-09	1 0F-03	131	0.36	7.4	None	1	1 OF-12	1308	F I	l 1.8F-1∩	768	SW	4 8F-10	
151	1318	FHE-26	Sample preparation	Pu-239 Am-241	1.0E-09 2.0E-10	1.0E-03 1.0E-03	13.1	0.36	7.4	None	1	1.0E-12 2.0E-13	1308	E	1.8E-10	768	SW	4.8E-10	2

Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated			ose Requirement		/y Monitoring		Source
					with Potential for	State	Height (m)	Diameter	Velocity	Device(s)	Abatement	Annual Emissions	Distance to			Distance	Direction	Unabated	Category
					Release (Ci)	Factor		(m)	(m/s)		Factor	(Ci)	SWMEI (m)	to SWMEI	(mrem)	to MEI (m)	to MEI	EDE (mrem)	
151	1322	FHE-33	Sample preparation	Gross alpha	6.0E-08	1.0E-03	12.8	0.36	8.1	None	1	6.0E-11	1308	E	1.2E-05	768	SW	3.1E-05	2
				Gross beta	1.0E-07	1.0E-03					·	1.0E-10							
				Gross gamma	1.0E-07	1.0E-03						1.0E-10							
				U-238	2.4E-04	1.0E-03						2.4E-07							
				U-235	3.1E-06	1.0E-03						3.1E-09							
				U-234	2.2E-05	1.0E-03						2.2E-08							
151	1326	FHE-42, 43, 44, 45	Radiochemical analysis of	Cf-249	1.0E-07	1.0E-03	12.8	0.36	6.8	None	1	1.0E-10	1308	E	2.9E-07	584	WNW	8.6E-07	1
		, -, , -	heavy element samples and	Pu-238	1.0E-07	1.0E-03						1.0E-10							
			weapon debris	Am-243	1.0E-07	1.0E-03						1.0E-10							
				Am-241	1.0E-07	1.0E-03						1.0E-10							
				Pu-236	1.0E-07 1.0E-07	1.0E-03 1.0E-03						1.0E-10 1.0E-10							
				Pu-239 Pu-240	1.0E-07 1.0E-07	1.0E-03						1.0E-10							
				Pu-242	1.0E-07	1.0E-03						1.0E-10							
				Pa-231	1.0E-07	1.0E-03						1.0E-10							
				Np-237	1.0E-07	1.0E-03						1.0E-10							
				U-238	1.0E-07	1.0E-03						1.0E-10							
	1330			U-235 U-234	1.0E-07 1.0E-07	1.0E-03 1.0E-03						1.0E-10 1.0E-10							
				Cm-246	1.0E-07	1.0E-03						1.0E-10							
				Cm-248	1.0E-07	1.0E-03						1.0E-10							
				Cs-137	1.0E-07	1.0E-03						1.0E-10							
				Sr-90	1.0E-07	1.0E-03						1.0E-10							
151		ELE SO	Touristant	0. 107	0.05.00	1.05.00	10.0	0.00	7.0			0.65.11	1000	-	1.25.00	700	6111	2.25.00	1
151		FHE-52	Transfer of waste samples for analysis	Cs-137 Co-60	9.6E-08 4.7E-09	1.0E-03 1.0E-03	12.8	0.36	7.6	None	1	9.6E-11 4.7E-12	1308	E	1.2E-09	768	SW	3.3E-09	2
			TOT attalysis	Sr-90	5.2E-08	1.0E-03						5.2E-11							
				Th-228	3.7E-12	1.0E-03						3.7E-15							
				Th-230	1.1E-11	1.0E-03						1.1E-14							
				Th-232	7.7E-13	1.0E-03						7.7E-16							
				Pu-238	1.1E-10	1.0E-03						1.1E-13							
				Pu-239	5.0E-09	1.0E-03						5.0E-12							
				Pu-240 Pu-241	3.0E-09 2.6E-09	1.0E-03 1.0E-03						3.0E-12 2.6E-12							
				Am-241	2.6E-10	1.0E-03						2.6E-13							
				U-234	7.4E-11	1.0E-03						7.4E-14							
				U-235	4.3E-12	1.0E-03						4.3E-15							
				U-238	1.3E-11	1.0E-03						1.3E-14							
				H-3	8.4E-12	1.0E-03						8.4E-15							
151	2103	FHE-6	Sorption studies	Pu-239	1.4E-07	1.0E-03	12.8	0.41	7.5	None	1	1.4E-10	1308	E	2.5E-08	768	SW	6.5E-08	2
131	2100	THE	Sorption studies	Pu-240	3.1E-08	1.0E-03	12.0	0.11	7.5	Hone		3.1E-11	1300		2.52 00	7.00	311	0.02 00	
				Pu-241	4.8E-07	1.0E-03						4.8E-10							
				Am-241	8.4E-09	1.0E-03						8.4E-12							
				Pu-238	4.0E-09	1.0E-03						4.0E-12							
151	2107	FHE-14	Transfer of solutions	Pu-239	2.0E-13	1.0E-03	12.8	0.41	7.3	None	1	2.0E-16	1308	E	3.8E-07	768	SW	9.8E-07	2
131	2107	FRE-14	for analysis	U-238	4.7E-06	1.0E-03 1.0E-03	12.0	0.41	1.3	NOTIE	1	4.7E-09	1300		3.0E-U/	100	SVV	3.0E-U/	
			Tot dilayou	U-235	2.2E-07	1.0E-03						2.2E-10		1					
				U-234	3.4E-06	1.0E-03						3.4E-09							
151	2109	FHE-19	Collection of daughter products	Th-228	1.2E-10	1.0E-06	13.1	0.30	6.1	None	1	1.2E-16	1308	E	1.0E-14	584	WNW	3.0E-14	2
			of Th-228								+			1		-			1
151	2109	FHE-15	Ion exchange studies	Sn-113	1.8E-08	1.0E-03	13.1	0.30	6.2	None	1	1.8E-11	1308	E	8.8E-13	584	WNW	2.5E-12	2
151	2117	FHE-23	Preparation of waste samples	Gross alpha	9.8E-09	1.0E-03	12.8	0.41	8.0	None	1	9.8E-12	1308	E	1.3E-09	768	SW	3.2E-09	2
			for analysis	Gross beta	2.2E-10	1.0E-03						2.2E-13		+					
151	2121	FHE-36	Sample preparation	Cs-137	9.5E-07	1.0E-03	12.8	0.41	8.0	None	1	9.5E-10	1308	E	1.5E-08	768	SW	3.8E-08	2
	=:=:	72 00	pro proparation	Co-60	4.7E-08	1.0E-03		1	5.5		·	4.7E-11	1.555		1.52.00	1 . 55		2.02.00	<u> </u>
				Sr-90	5.1E-07	1.0E-03						5.1E-10							
				Th-228	3.7E-11	1.0E-03						3.7E-14							
				Th-230	1.1E-10	1.0E-03						1.1E-13		1					1
			-	Th-232 Pu-238	7.6E-12 1.1E-09	1.0E-03 1.0E-03						7.6E-15 1.1E-12		+	-				
			+	Pu-238 Pu-239	4.9E-08	1.0E-03 1.0E-03						4.9E-11							
			1	Pu-240	2.9E-08	1.0E-03						2.9E-11		1					
				Pu-241	2.6E-08	1.0E-03						2.6E-11							
				Am-241	2.6E-09	1.0E-03						2.6E-12							
				U-234	7.3E-10	1.0E-03						7.3E-13							

Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated	10 mrem/y	Site-Wide Do	se Requirement	0.1 mrem	y Monitoring	Requirement	Source
J			·		with Potential for	State	Height (m)	Diameter	Velocity	Device(s)	Abatement	Annual Emissions	Distance to		EDE	Distance	Direction	Unabated	Category
					Release (Ci)	Factor		(m)	(m/s)		Factor	(Ci)	SWMEI (m)	to SWMEI	(mrem)	to MEI (m)	to MEI	EDE (mrem)	
151	2121	(continued)		U-235	4.3E-11	1.0E-03						4.3E-14							
				U-238	1.3E-10	1.0E-03						1.3E-13							
				Pu-239	2.4E-08	1.0E-03						2.4E-11							+
			+	Sr-90 H-3	4.9E-10 7.3E-08	1.0E-03 1.0E-03						4.9E-13 7.3E-11							+
				11-3	7.3L-00	1.0L-03						7.3L-11				+			+
151	2131	FHE-56	Transfer and processing of	Pu-239	2.0E-11	1.0E-06	12.8	0.41	7.3	None	1	2.0E-17	1308	E	3.7E-15	584	WNW	8.7E-15	1
	2.0.	FHE-47	glass samples for	Pu-238	3.0E-12	1.0E-06	12.8	0.41	7.8	None	1	3.0E-18	1000		0.7.2.10			0.1.2.10	+
			radiochemical analysis	U-235	3.0E-12	1.0E-06						3.0E-18							
				U-238	3.0E-12	1.0E-06						3.0E-18							
				Cs-137	6.0E-11	1.0E-06						6.0E-17							
				Sr-90	6.0E-11	1.0E-06						6.0E-17							
				Eu-152	6.0E-12	1.0E-06						6.0E-18							
				Eu-154	6.0E-12	1.0E-06						6.0E-18							
151	2122	EUE	6	0	1.45.14	1.05.02	12.0	0.41	0.1	Maria	1	1 45 17	1200	-	1.05.15	700	CIA	4 55 45	
151	2133	FHE-57	Swipe sample analysis	Gross alpha	1.4E-14	1.0E-03	12.8	0.41	8.1	None	1	1.4E-17	1308	E	1.8E-15	768	SW	4.5E-15	2
151	2143	FHE-63	Transfer of standards for	H-3	3.6E-12	1.0E-03	12.8	0.41	8.2	None	1	3.6E-15	1308	E	8.7E-19	768	SW	2.1E-18	2
131	2143	FHE-03	the analysis of environmental	п-э	3.0E-12	1.0E-03	12.0	0.41	0.2	None	ı	3.0E-13	1306		0.76-19	700	SVV	2.15-10	+ -
			samples; analysis of standards																+
			for environmental samples																+
151	2147	FHE-67	Transfer of yield tracers for	Pu-242	7.1E-13	1.0E-03	12.8	0.41	8.0	None	1	7.1E-16	1308	E	6.5E-14	768	SW	1.6E-13	2
151	2149	FHE-78	Transfer of yield tracers	Pu-238	2.0E-14	1.0E-03	13.1	0.41	7.8	None	1	2.0E-17	1308	E	6.0E-13	768	SW	1.5E-12	2
	51 2149 FHE-78		samples as yield tracers	Pu-239	4.0E-14	1.0E-03						4.0E-17							
			during analysis	Pu-240	4.0E-14	1.0E-03						4.0E-17							
				Pu-242	3.0E-12	1.0E-03						3.0E-15							
				U-232	1.0E-12	1.0E-03						1.0E-15				-			
				U-233	9.0E-13	1.0E-03						9.0E-16							
				U-238	4.0E-15 1.4E-12	1.0E-03 1.0E-03						4.0E-18							+
				Cs-134 Cs-137	8.1E-13	1.0E-03 1.0E-03						1.4E-15 8.1E-16							+
				C3-137	0.1L-13	1.0L-03						0.1L-10				+			+
151	2302A	FHE-9	Waste treatability studies	H-3	1.0E-04	1.0E-03	13.1	0.41	7.5	None	1	1.0E-07	1308	E	2.1E-11	768	SW	5.9E-11	2
101	2002/1	1112 0	Tracto di catability ditadico	U-235	6.1E-14	1.0E-03		0	1.0	110110		6.1E-17	1.000				0	0.02	+
				0 000								, , , , , , , , , , , , , , , , , , ,							+
151	2308	FHE-16	Ceramics leaching studies	Pu-239	1.9E-02	1.0E-03	12.8	0.41	7.3	Double HEPA	0.0001	1.9E-09	1308	Е	3.5E-07	768	SW	8.9E-03	2
				Pu-240	4.2E-03	1.0E-03						4.2E-10							
				Pu-241	6.8E-02	1.0E-03						6.8E-09							
				Am-241	1.1E-03	1.0E-03						1.1E-10							
				Pu-238	5.6E-04	1.0E-03						5.6E-11							
				U-234	2.2E-07	1.0E-03						2.2E-14							
				U-235	9.7E-09	1.0E-03						9.7E-16				-			+
				U-238	2.1E-07	1.0E-03						2.1E-14							+
151	2308	FHE-12	Ceramics leaching studies	U-234	6.6E-07	1.0E-03	13.1	0.41	7.8	None	1	6.6E-10	1308	E	5.9E-08	768	SW	1.4E-07	2
131	2300	1112.12	ceramics leaching studies	U-235	2.9E-08	1.0E-03	13.1	0.41	7.0	None	'	2.9E-11	1300		3.32 00	700	3**	1.42 07	+ -
				U-238	6.1E-07	1.0E-03						6.1E-10							+
151	2312	FHE-21	Solubility studies	Np-237	2.7E-08	1.0E-03	12.8	0.41	7.6	Double HEPA	0.0001	2.7E-15	1308	E	4.9E-13	768	SW	1.2E-08	2
151	2312	FHE-21	Solubility studies	Np-237	6.4E-09	1.0E-03	12.8	0.41	7.1	None	1	6.4E-12	1308	E	1.2E-09	768	SW	3.0E-09	2
151	2318	FHE-22	Transfer of sample solutions	Pu-242	1.9E-09	1.0E-03	9.8	0.41	8.0	Double HEPA	0.0001	1.9E-16	1308	E	2.5E-14	768	SW	7.8E-10	2
151	2322	FHE-38	Transfer and processing of	Cs-137	3.4E-07	1.0E-03	12.8	0.41	8.1	None	1	3.4E-10	1308	Е	5.6E-09	768	SW	1.4E-08	2
131	4344	FПE-38	of waste sludge samples	Cs-137 Co-60	3.4E-07 1.7E-08	1.0E-03 1.0E-03	12.8	0.41	0.1	inorie	1	3.4E-10 1.7E-11	1308	E	3.0E-U9	700	SVV	1.46-08	+ 4
			for TCLP, STLC, pH, % moisture,	Sr-90	1.8E-07	1.0E-03						1.8E-10				+			+
			TTLC analyses	Th-228	1.3E-11	1.0E-03						1.3E-14							+
			1120 unalyoco	Th-230	3.9E-11	1.0E-03						3.9E-14							+
				Th-232	2.8E-12	1.0E-03						2.8E-15							
				Pu-238	3.9E-10	1.0E-03						3.9E-13							
				Pu-239	1.8E-08	1.0E-03						1.8E-11							
		<u> </u>		Pu-240	1.1E-08	1.0E-03						1.1E-11							
		·		Pu-241	9.4E-09	1.0E-03						9.4E-12						<u> </u>	
				Am-241	9.4E-10	1.0E-03						9.4E-13							
				U-234	2.6E-10	1.0E-03			1	1		2.6E-13	-			-			
				U-235	1.5E-11	1.0E-03			1			1.5E-14							
				U-238	4.7E-11	1.0E-03						4.7E-14	-						+
				Gross alpha	9.0E-09	1.0E-03 1.0E-03				-		9.0E-12 2.1E-13	-	+					+
				Gross beta H-3	2.1E-10 2.1E-08	1.0E-03 1.0E-03						2.1E-13 2.1E-11	 			+			+
				п-3	∠.1E-Uδ	1.UE-U3	l	I	1			Z.IE-II	l	1					

Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated	10 mrem/y S	Site-Wide Do	se Requirement	0.1 mrem/	y Monitoring I	Requirement	Source
			·		with Potential for	State	Height (m)	Diameter	Velocity	Device(s)	Abatement	Annual Emissions	Distance to		EDE	Distance	Direction	Unabated	Category
					Release (Ci)	Factor		(m)	(m/s)		Factor	(Ci)	SWMEI (m)	to SWMEI	(mrem)	to MEI (m)	to MEI	EDE (mrem)	
151	2326	FHE-39	Chamical analysis of wasta	Cs-137	3.8E-07	1.0E+00	12.8	0.41	7.6	None	1	3.8E-07	1308	E	8.8E-06	768	SW	2.2E-05	2
151	2320	LUE-23	Chemical analysis of waste	Co-60	1.8E-08	1.0E+00	12.0	0.41	7.0	None	1	1.8E-08	1306	<u> </u>	0.0E-00	700	SW	2.ZE-U3	
				Sr-90	2.0E-07	1.0E+00						2.0E-07							
				Th-228	1.5E-11	1.0E+00						1.5E-11							
				Th-230	4.3E-11	1.0E+00						4.3E-11							
				Th-232	3.0E-12	1.0E+00						3.0E-12							
				Pu-238	4.4E-10	1.0E+00						4.4E-10							
				Pu-239	2.0E-08	1.0E+00						2.0E-08							
				Pu-240	1.2E-08	1.0E+00						1.2E-08							
				Pu-241	1.0E-08	1.0E+00						1.0E-08							
				Am-241	1.0E-09	1.0E+00						1.0E-09							
				U-234 U-235	2.9E-10 1.7E-11	1.0E+00 1.0E+00						2.9E-10 1.7E-11							
				U-238	5.2E-11	1.0E+00						5.2E-11							+
				Gross alpha	3.0E-08	1.0E+00						3.0E-08							
				Gross beta	6.0E-10	1.0E+00						6.0E-10							
				H-3	9.0E-08	1.0E+00						9.0E-08							
151	2326A	FHE-40	Preparation of environmental	Gross alpha	1.0E-09	1.0E-03	12.8	0.30	3.9	None	1	1.0E-12	1308	E	1.4E-10	584	WNW	5.4E-10	2
			and waste samples																
151	2226	ELE EA	Andring	D 000	2.05.00	1.05.00	10.0	0.11	7.5			2.05.12	1000	-	F 15 10	700	6)**	1 25 22	-
151	2330	FHE-50	Analysis of standards for	Pu-239	3.9E-09	1.0E-03	12.8	0.41	7.5	None	1	3.9E-12	1308	E	5.1E-10	768	SW	1.3E-09	2
			waste samples; analysis	H-3 H-3	5.7E-12 9.0E-09	1.0E-03 1.0E+00						5.7E-15 9.0E-09	-						
			of waste samples	H=3	3.UE-U3	1.02+00						3.UE-U3	+			1	 		
151	2344	FHE-65	Preparation of organic liquid	H-3	1.0E-09	1.0E-03	12.8	1.33	7.9	None	1	1.0E-12	1308	E	3.0E-11	1469	NE	1.2E-10	1
		FHE-69	samples for VOC analysis	C-14	1.0E-11	1.0E-03	12.8	1.33	8.0	None	1	1.0E-14	1000	_	0.02				
			by GC/MS	Sr-90	1.0E-10	1.0E-03						1.0E-13							
			•	Cs-137	1.0E-10	1.0E-03						1.0E-13							
				Eu-152	1.0E-11	1.0E-03						1.0E-14							
				U-238	1.0E-09	1.0E-03						1.0E-12							
				U-235	1.0E-10	1.0E-03						1.0E-13							
				Pu-239	1.0E-10	1.0E-03						1.0E-13							
				Pu-238	1.0E-10	1.0E-03						1.0E-13							
-				Am-241	1.0E-10	1.0E-03						1.0E-13							
151	2348	FHE-75	Freeze trapping/analysis of tritium	H-3	1.5E-13	1.0E+00	12.8	0.41	8.7	None	1	1.5E-13	1308	E	3.7E-17	768	SW	8.7E-17	2
131	23 10	THE 75	Treeze trapping/ analysis of tritain	H-3	3.6E-12	1.0E-03	12.0	0.11	0.1	None	· ·	3.6E-15	1300	_	3.72.17	7.00		0.72 17	
151	2350	FHE-76	Transfer of tracer solutions	Pu-242	1.2E-12	1.0E-03	12.8	0.41	8.4	None	1	1.2E-15	1308	E	4.6E-13	768	SW	1.1E-12	2
				Am-243	5.7E-13	1.0E-03						5.7E-16							
				U-232	3.3E-13	1.0E-03						3.3E-16							
				Pu-239	7.3E-13	1.0E-03						7.3E-16							
				Am-241	8.6E-14	1.0E-03						8.6E-17							
				U-234	6.8E-11	1.0E-03						6.8E-14				1			
				U-235	8.5E-09	1.0E-03						8.5E-12	-			1			1
				U-238	1.2E-06	1.0E-03					_	1.2E-09	-			1			1
Buildings 175		rt of the Uranium Atomic Va	 apor Laser Isotope Separation (U-AVLIS) p	rogram, affiliated w	uith The United States	Enrichment (Corporation (USF)	L C). In June 1999	. USEC suspender	d further developm	ent of the II-AVLIS t	echnology							
			ning in early 2002, and the sampling syste		The shired states				, 1010 Suspender	acreiopin									
*Gross alpha	and Gross beta em	nissions are continuously mo	onitored at the stack. Monitoring data, rath	ner than the invento															
**Because me	onitoring takes pla	ce after HEPA filtration, an	unabated EDE cannot be determined (see	discussion in Sectio	n II, subsection "Stac	k Monitoring fo	or Gross Alpha ar	nd Gross Beta Rad	diation.")										
					ļ							0.55.65	ļ			ļ			1 -
175	103	FFE-02	Operations discontinued	Gross alpha	*	NA	9.4	0.61	4.5	HEPA	1.0E-02	0.0E+00	**	**	0.0E+00	**	**	**	3
	103	FFE-01		Gross beta	*	NA	9.4	0.61	4.6			0.0E+00	-						1
	112 112	FHE-02 FHE-01			-		6.8 6.7	0.36	6.4				-			1			+
	128	FHE-2000			+		8.9	0.59	4.6				 						1
	128	FHE-1000			<u> </u>		8.9	0.59	5.2				+						
		111E 1000					0.5	0.55	5.2										
177	1020	FHE-22	Operations discontinued	Gross alpha	*	NA	6.4	0.30	8.9	HEPA	0.01	0.0E+00	**	**	0.0E+00	**	**	**	3
				Gross beta	*	NA						0.0E+00							
			Space Technology Directorate. The facility	houses a high-ener	gy linear accelerator	(LINAC) and re	esearch laborator	ies.											
The accelerat	tor beam can produ	uce small quantities of short	t-lived air activation products.										-			1			
194	D122	TE-FE4	Lingo goodgrater voult	0-15	6.0E-02	1.0E+00	30.5	1.37	4.5	None	1	6.0E-02	1525	SSE	E 2F 07	E20	NE	E 2F 0F	1 2
134	B122	(Target Exhaust)	Linac accelerator vault	N-13	1.1E-01	1.0E+00 1.0E+00	30.3	1.37	4.5	None	1	1.1E-01	1323	SSE	5.2E-07	538	INE	5.3E-05	2
		(Talyet Exilaust)		IN-13	1.15-01	1.02+00						1.16-01	-						
					1														

Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated		ite-Wide Do	se Requirement		y Monitoring		Source
					with Potential for	State	Height (m)	Diameter	Velocity	Device(s)		Annual Emissions	Distance to	Direction	EDE	Distance	Direction	Unabated	Category
					Release (Ci)	Factor		(m)	(m/s)		Factor	(Ci)	SWMEI (m)	to SWMEI	(mrem)	to MEI (m)	to MEI	EDE (mrem)	
194	B124	TE-FE4	Storage	Na-22	5.0E-05	1.0E-03	30.5	1.37	4.5	None	1	5.0E-08	1525	SSE	9.7E-09	538	NE	1.4E-07	2
				U-233	1.2E-05	1.0E-06						1.2E-11					<u> </u>		
				U-234	6.0E-09	1.0E-06						6.0E-15					<u> </u>		
				U-235	3.9E-06	1.0E-06						3.9E-12					<u> </u>		
				U-236	3.5E-08	1.0E-06						3.5E-14					<u> </u>		
				U-238	1.0E-04	1.0E-06						1.0E-10					<u> </u>		
194	B130	TE-FE4	Positron beam generation	0-15	5.5E-01	1.0E+00	30.5	1.37	4.5	None	1	5.5E-01	1525	SSE	5.2E-06	538	NE	5.2E-04	2
131	B130	12121	1 osition beam generation	N-13	1.1E+00	1.0E+00	30.3	1.07	1.5	None		1.1E+00	1020	332	3.22 00	330		5.22 01	
194	1131	Room Air	Positron materials science	Na-22	3.6E-06	1.0E-03	NA	NA	NA	None	1	3.6E-09	1525	ESE	2.4E-09	532	W	7.0E-08	2
			experiments														 '		
D '' I' 010				10: 5: .													 '		
			ology Directorate (formerly the Physica from past operations of the rotating ta		·		ments.										<u> </u>		
The current	radionuciide emissio	ons are due to contamination	Trom past operations of the rotating ta	Teutron source	, which is no longer in	operation.											 '		
212	174	FHE-7	Contamination	H-3	1.7E-02	1.0E-06	4.3	0.5	0.5	None	1	1.7E-08	1278	ENE	8.5E-12	38	SW	3.4E-10	2
	.,,	11121	Contamination	11.5	1.72 02	1.02 00	1.5	0.3	0.5	None		1.72 00	1270	LIVE	0.52 12	30		3.12.10	
212	184	Room Air	Contamination	H-3	1.0E-03	1.0E-06	NA	NA	NA	None	1	1.0E-09	1278	ENE	5.0E-13	38	SW	2.2E-11	2
The 231 cor	mplex houses resear	rch and development activities	s conducted by the Chemistry and Mate	erials Science Directo	orate, Engineering, We	apons Engine	ering, and Safegu	ards and Security	/ Materials Manag	ement Division.									
Management	t oversight for Buildi	ling 231 is provided by the En	gineering Directorate through the Engir	neering Sciences Divi	ision.														
		·								·					·				
231	1000	FFE-5	Metal casting	U-238	2.6E-07	1.0E-06	8.2	0.32	7.7	HEPA	0.01	2.6E-15	1167	E	1.7E-11	671	W	6.8E-11	2
				U-235	3.3E-09	1.0E-06						3.3E-17					L		
				U-234	2.4E-08	1.0E-06						2.4E-16					<u> </u>		
																	<u> </u>		
231	1427	Room Air	Wet grinding/lapping	U-238	3.1E-07	1.0E-06	NA	NA	NA	None	1	3.1E-13	1167	E	4.3E-11	671	W	3.6E-10	2
				U-235	1.5E-08	1.0E-06						1.5E-14					 '		
				U-234	3.3E-07	1.0E-06						3.3E-13							
221	1600	D A'.	Edular Landau and Pal	11 220	1.55.00	1.05.00	N/A		210	Nicos	1	1.55.00	1167	_	1 15 07	671	<u> </u>	0.05.07	
231	1600	Room Air	Friction test on solid	U-238	1.5E-03	1.0E-06	NA	NA	NA	None	1	1.5E-09	1167	E	1.1E-07	671	W	8.8E-07	2
			depleted uranium bars	U-235	1.9E-02 1.4E-01	1.0E-06						1.9E-08 1.4E-07					 		
				U-234	1.4E-01	1.0E-06						1.4E-07							
231	1640	Room Air	Mechanical test; quasistatic	U-238	5.9E-09	1.0E-06	NA	NA	NA	None	1	5.9E-15	1167	E	4.1E-13	671	W	3.4E-12	2
231	1040	ROOM All	compression	U-235	7.6E-11	1.0E-06	INA	INA	INA	None	1	7.6E-17	1107		T.1L-13	071		J.4L-12	
			compression	U-234	5.5E-10	1.0E-06						5.5E-16							
				0 20 .	0.02 . 0	1102 00						0.02 . 0							
231	1678	Room Air	Mechanical test; compression	U-238	6.8E-09	1.0E-06	NA	NA	NA	None	1	6.8E-15	1167	Е	4.7E-13	671	W	3.9E-12	2
			Hopkinson bar (U6Nb)	U-235	8.7E-11	1.0E-06						8.7E-17	-						
				U-234	6.3E-10	1.0E-06						6.3E-16							
231	1737	FGBE-5	Electron beam welding	U-238	1.5E-06	1.0E-06	10.1	0.46	1.5	HEPA	0.01	1.5E-14	1167	E	1.0E-12	671	W	5.6E-10	2
				U-235	1.9E-08	1.0E-06						1.9E-16					<u> </u>		
				U-234	1.4E-07	1.0E-06						1.4E-15					<u> </u>		
				U-238	8.1E-11	1.0E-03						8.1E-16					<u> </u>		
				U-235	1.0E-12	1.0E-03						1.0E-17					<u> </u>		
				U-234	7.5E-12	1.0E-03						7.5E-17					 '		
221	17274	FUE E4	Flootron has a subdise	11 220	1.55.00	1.05.00	10.1	0.40	1.5	LIEDA	0.01	1 55 14	1107		1.05.13	671	14/	F 65 10	-
231	1737A	FHE-54	Electron beam welding	U-238	1.5E-06	1.0E-06	10.1	0.46	1.5	HEPA	0.01	1.5E-14	1167	E	1.0E-12	671	W	5.6E-10	2
				U-235 U-234	1.9E-08 1.4E-07	1.0E-06 1.0E-06		-				1.9E-16 1.4E-15					<u> </u>		
\vdash				U-234 U-238	8.1E-11	1.0E-06 1.0E-03		-			+	8.1E-16							
 				U-235	1.0E-12	1.0E-03 1.0E-03		1			+	1.0E-17							
				U-234	7.5E-12	1.0E-03					+	7.5E-17							
				0 20-7	1.02.12														
231	1737B	FHE-54	Electron beam welding	U-238	1.7E-07	1.0E-03	10.1	0.46	11.5	HEPA	0.01	1.7E-12	1167	E	1.1E-10	671	W	6.0E-08	2
	-			U-235	2.2E-09	1.0E-03						2.2E-14							T -
				U-234	1.6E-08	1.0E-03						1.6E-13							
231	1739	FGBE-5	Storage	U-238	1.5E-07	1.0E-06	10.1	0.46	11.5	HEPA	0.01	1.5E-15	1167	E	1.0E-13	671	W	5.4E-11	2
				U-235	2.0E-09	1.0E-06						2.0E-17							
				U-234	1.4E-08	1.0E-06						1.4E-16					<u> </u>		
																	<u> </u>		
231	1900HB	FGBE-7/8	Storage	U-238	4.9E-06	1.0E-06	2.4	0.20	14.4	None	1	4.9E-12	1167	E	3.3E-10	671	W	2.2E-09	2
				U-235	6.3E-08	1.0E-06		-				6.3E-14					 '		
				U-234	3.0E-07	1.0E-06		-				3.0E-13					 '		
	10444	Danie Ate	Machanitation	11 222	1 25 07	1.05.00	N/A	N14	N/A	NI	1	1 25 12	1107		0.25.12	671	14/	7.05.11	+ -
221		Room Air	Mechanical testing	U-238	1.3E-07	1.0E-06	NA	NA	NA	None	1	1.3E-13	1167	E	9.2E-12	671	W	7.6E-11	2
231	1944A			11 225															
231	1944A			U-235 U-234	1.7E-09 1.3E-08	1.0E-06 1.0E-06						1.7E-15 1.3E-14					<u> </u>		

Buildina	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated	10 mrom/v	Sita Wida Da	se Requirement	0.1 mrom	/v Monitorino	Requirement	Source
Building	ROOM/ Alea	Stack ID	Operation	Radionuclides	with Potential for	State	Height (m)	Diameter	Velocity	Device(s)	Abatement	Annual Emissions			EDE	Distance	Direction	Unabated	Category
					Release (Ci)	Factor		(m)	(m/s)		Factor	(Ci)	SWMEI (m)		(mrem)	to MEI (m)	to MEI	EDE (mrem)	
231	1945	FHE-40	Metal characterization	U-238	2.0E-09	1.0E-06	10.7	0.36	3.8	None	1	2.0E-15	1167	E	1.3E-13	671	W	5.4E-13	2
				U-235 U-234	2.6E-11 1.9E-10	1.0E-06 1.0E-06						2.6E-17 1.9E-16							+
				0 23 1	1.52 10	1.02.00						1.32 10							1
231	1945A	Room Air	Metal characterization	U-238	2.0E-12	1.0E-06	NA	NA	NA	None	1	2.0E-18	1167	E	1.4E-16	671	W	1.2E-15	2
				U-235 U-234	2.6E-14 1.9E-13	1.0E-06 1.0E-06						2.6E-20 1.9E-19							+
				0-234	1.9E-13	1.0E-06	+					1.9E-19							+
231	1945B	FHE-40	Metal characterization	U-238	1.4E-09	1.0E-03	10.0	0.41	4.6	None	1	1.4E-12	1167	E	8.9E-11	671	W	3.5E-10	2
				U-235	1.7E-11	1.0E-03						1.7E-14							
				U-234	1.3E-10	1.0E-03	+					1.3E-13						-	
231	1945C	Room Air	Metal characterization	U-238	2.0E-12	1.0E-06	NA	NA	NA	None	1	2.0E-18	1167	E	1.4E-16	671	W	1.2E-15	2
				U-235	2.6E-14	1.0E-06						2.6E-20							
				U-234	1.9E-13	1.0E-06						1.9E-19							
231	1945D	Room Air	Metal polishing	U-238	2.0E-09	1.0E-06	NA	NA	NA	None	1	2.0E-15	1167	E	1.4E-13	671	W	1.2E-12	2
231	13430	KOOTII AII	Metal polishing	U-235	2.6E-11	1.0E-06	INA	IVA	IVA	None	'	2.6E-17	1107		1.46-13	071	***	1.2L-12	
				U-234	1.9E-10	1.0E-06						1.9E-16							
221	10455	Dan Att.	Man mindle Co-Palitie	11 222	2.05.00	1.05.00	NIA.	N/A	N1 A	Ninn	1	2.05.00	1107	-	1 45 07	671	14/	1.25.00	+ -
231	1945E	Room Air	Wet grinding/polishing	U-238 U-235	2.0E-06 2.6E-08	1.0E-03 1.0E-03	NA	NA	NA	None	1	2.0E-09 2.6E-11	1167	E	1.4E-07	671	W	1.2E-06	2
				U-234	1.9E-07	1.0E-03						1.9E-10							\pm
									L										I
			irectorate. Operations in the facility in eterization studies; some is used for ic			surface, and	subsurface; precis	sion cutting, ion i	mplanting, and r	metallurgical studies.	1								+
MOST OF THE	e depieted draillum	i iii uiis bullullig is useu for charac	Studies, Some is used for it	n beam impiantation	experiments.		+											 	+
235	1122	FHE-1A/1B, FHE2A/2B,	Surface analysis	U-234	1.5E-11	1.0E-06	10.7	2.75	4.0	None	1	1.5E-17	1065	ENE	1.3E-14	556	SW	1.3E-14	2
		FGBE-1A/1B		U-235	2.1E-12	1.0E-06						2.1E-18							
				U-238	1.6E-10	1.0E-06						1.6E-16							+
*Gross alph	na and Gross beta	emissions are continuously monitories.	Dred at the stack. Monitoring data, rat	l her than the invento	ry approach, are used	l d to determin	ne emissions.												+
**Because	monitoring takes p	place after HEPA filtration, an una	bated EDE cannot be determined (see	discussion in Section	n II, subsection "Stac	k Monitoring	for Gross Alpha a	nd Gross Beta Ra	diation.")										
225	1120	FUE 1 A /1 D FUE 2 A /2 D	Decreasion of aluterium	Curan alala	*	NIA	10.7	2.75	4.0	Davida UEDA	0.0001	0.05.00	**	**	0.05.00	**	**	**	+ -
235	1130	FHE-1A/1B, FHE2A/2B, FGBE-1A/1B	Preparation of plutonium samples for diamond anvil studies	Gross alpha Gross beta	*	NA NA	10.7	2.75	4.0	Double HEPA	0.0001	0.0E+00 0.0E+00	**	^^	0.0E+00	^^	^^	^^	3
		TODE 170 TO	Samples for diamona arryll scales	Gross Beta		10/						0.02100							
235	1131	HDCH-6,7	Metallographic sample	U-234	1.1E-08	1.0E-06	10.7	2.75	4.0	HEPA	0.01	1.1E-16	1065	ENE	9.2E-14	556	SW	9.6E-12	2
		(FHE-1A/1B, FHE2A/2B, FGBE-1A/1B)	preparation	U-235 U-238	1.5E-09 1.2E-07	1.0E-06 1.0E-06						1.5E-17 1.2E-15							+
		FGBE-TA/TB)		0-236	1.25-07	1.0E-06						1.2E-13							+
235	1133	FHE-1A/1B, FHE2A/2B,	Microstructure examination	U-234	2.8E-09	1.0E-06	10.7	2.75	4.0	None	1	2.8E-15	1065	ENE	2.3E-12	556	SW	2.4E-12	2
		FGBE-1A/1B		U-235	3.8E-10	1.0E-06						3.8E-16							
				U-238	3.0E-08	1.0E-06						3.0E-14							+
235	1235	FHE-1A/1B, FHE2A/2B,	X-ray diffraction of	U-234	1.1E-09	1.0E-06	10.7	2.75	14.3	None	1	1.1E-15	1065	ENE	1.5E-13	556	SW	1.6E-13	2
		FGBE-1A/1B	uranium oxide ceramics	U-235	4.6E-11	1.0E-06						4.6E-17							
				U-238	9.9E-10	1.0E-06						9.9E-16							+
Building 24	11 is administered I	by the Chemistry and Material Sci	l ences Directorate for material propert	l ties research and tes	l ting, and for study of	soil bacteria													+
241	1616	Room Air	Paritcle size analysis of powders	U-238	2.0E-11	1.0E-03	NA	NA	NA	None	1	2.0E-14	1140	E	2.9E-12	697	W	2.2E-11	2
				U-235 U-234	9.3E-13 2.1E-11	1.0E-03 1.0E-03	+					9.3E-16 2.1E-14							+
241	1678	FHE-55	Research and development	U-238	1.4E-05	1.0E+00	7.9	0.28	15.4	HEPA	0.01	1.4E-07	1140	E	1.8E-05	821	SW	4.4E-03	2
			of methods for radionuclide	U-235	6.7E-07 1.5E-05	1.0E+00	-					6.7E-09 1.5E-07						-	
			immobilization using uranium oxide	U-234 U-238	9.4E-07	1.0E+00 1.0E-03	+					9.4E-12	-					-	+
				U-235	4.4E-08	1.0E-03						4.4E-13							
				U-234	1.0E-06	1.0E-03						1.0E-11							
				U-238 U-235	1.9E-07 8.8E-09	1.0E-06 1.0E-06	+					1.9E-15 8.8E-17						-	+
				U-234	2.0E-07	1.0E-06						2.0E-15							+
241	1838	FGBE-10	Pressing and sintering of	U-238	8.9E-07	1.0E+00	7.6	0.15	12.9	HEPA	0.01	8.9E-09	1140	E	1.2E-06	697	W	5.9E-04	2
			uranium oxide disks	U-235 U-234	4.2E-08 9.6E-01	1.0E+00 1.0E+00	+					4.2E-10 9.6E-03							+
				U-238	2.0E-10	1.0E-03						2.0E-15							
				U-235	9.5E-12	1.0E-03						9.5E-17							I
				U-234 U-238	2.2E-10 9.9E-15	1.0E-03 1.0E-06	-	-			1	2.2E-15 9.9E-23						-	+
				U-235	9.9E-15 4.6E-16	1.0E-06 1.0E-06	+					9.9E-23 4.6E-24			1			 	+
				U-234	1.1E-14	1.0E-06						1.1E-22							

Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated			se Requirement	0.1 mrem/	y Monitoring		Source
					with Potential for	State	Height (m)	Diameter	Velocity	Device(s)	Abatement	Annual Emissions	Distance to		EDE	Distance	Direction	Unabated	Category
					Release (Ci)	Factor		(m)	(m/s)		Factor	(Ci)	SWMEI (m)	to SWMEI	(mrem)	to MEI (m)	to MEI	EDE (mrem)	
														_					
241	1838	FHE-7	Weighing and measuring of	U-238	4.6E-07	1.0E+00	7.9	0.39	6.6	None	0.01	4.6E-09	1140	E	6.4E-07	697	W	2.5E-04	2
			sintered uranium oxide disks	U-235	2.2E-08	1.0E+00						2.2E-10							
				U-234	5.0E-07	1.0E+00						5.0E-09							
				U-238	9.9E-09	1.0E-03						9.9E-14							
				U-235	4.6E-10	1.0E-03						4.6E-15							
				U-234	1.1E-08	1.0E-03						1.1E-13							
2.41	1041	EUE EO	Control of books in the control	0.14	2.05.07	1.05.00	7.0	0.20	11.0	Maria	1	2.05.07	1140	-	4.25.00	607	14/	1.25.00	+
241	1841	FHE-53	Study of bacterial conversion	C-14	2.0E-07	1.0E+00	7.9	0.30	11.3	None	1	2.0E-07	1140	E	4.2E-09	697	W	1.3E-08	2
			of organic carbon in waste	C-14	4.3E-10	1.0E-03						4.3E-13				754	WNW	1.3E-08	
			to carbon dioxide																
241	1000	D A.	THE SAPERSON AND AREA SALE	D 22	0.25.00	1.05.02			N14	Maria	1	6.25.11	1110	-	1 55 12	754	14/5 DA/	1.05.11	+ -
241	1886	Room Air	Hybridization studies with	P-32	6.3E-08	1.0E-03	NA	NA	NA	None	1	6.3E-11	1140	E	1.5E-12	754	WNW	1.0E-11	2
			nucleic acids from soil bacteria																
D 11 11 01	54 1 11 51		0 () 0 () 15 () 18	5				9											+
			Safety, Security and Environmental Pro							are stored until they	can be disposed.								
			ge from earthquakes. Room exhausts fro		ea are double HEPA fil	tered; glove b	ox exhausts are t	riple HEPA filtere	ed.										
			continuously sampled by simple filter sy																
			PA/DOE Memorandum of Understanding																
**Because	monitoring takes pla	lace after HEPA filtration, an ur	nabated EDE cannot be determined (see	discussion in Section	on II, subsection "Stac	k Monitoring f	for Gross Alpha ar	nd Gross Beta Ra	diation.")										
	Unhardened Area*																		
251	1003	FHE-5	General chemistry	Gross alpha	*	NA	4.3	0.26	8.6	HEPA	0.01	0.0E+00	1188	E	0.0E+00	**	**	**	3
	1003	FHE-4		Gross beta			4.3	0.27	4.2			0.0E+00							1
	1142	FHE-8					4.3	0.32	4.1										T
	1142	FHE-9					4.3	0.26	5.1				1						
	1142	FHE-10					4.3	0.28	13.7										1
	1150	FGBE-33,34					8.0	0.15	12.8										+
	1150	FFE-15					4.3	0.31	7.6										+
	1165	FGBE-31,32					5.5	0.87	0.1										+
	1211	FHE-6					6.4	0.25	8.0										+
	1211	FHE-7					6.4	0.25	4.3										+
	1212	FGBE-15,16					5.5	0.10	8.0										+
	1232	FGBE-38,39					7.2	0.15	5.1										+
	1232						-												
		FFE-9					4.3	0.19	14.7										
	1235	FFE-12					4.3	0.25	7.6										
	1235	FGBE-29,30					5.5	0.13	7.1										
	1363	FHE-12					4.3	0.32	9.1										
	1363	FHE-13					6.4	0.28	6.8										
	1364	FFE-23					4.3	0.34	9.1										
	1364	FGBE-35,36					4.3	0.13	11.2										
	1314, 1354	FGBE-44,45					10.2	0.15	10.2										
	Hot cells	FGBE-40,41					5.5	0.23	5.6										
	Hot cells	FGBE-42,43					5.5	0.36	12.7										
	1150	FFE-13					5.5	0.28	4.1										
																			1
	Hardened Area																		
251	Glove Boxes*	FGBE-1000	Previous transuranic research	Gross alpha	*	NA	7.8	0.30	4.8	Triple HEPA	0.000001	0.0E+00	1188	E	0.0E+00	**	**	**	3
		FGBE-2000		Gross beta			7.8	0.30	4.8			0.0E+00							
	Room Exhaust*	FFE-1000		Gross alpha	*	NA	7.8	0.50	11.7	Double HEPA	0.0001	0.0E+00	1188	E	0.0E+00	**	**	**	3
		FFE-2000		Gross beta			7.8	0.50	11.7			0.0E+00							
Buildina 2	53 houses the Hazar	rds Control Department, and th	ne facility includes laboratories for the o	hemical analysis and	l counting of radioacti	ve samples.													1
3		.,		,	J : 2:2::2:10t.	1													T
253	1708	Room Air	Gross alpha/beta analysis of	Pu-238	2.1E-10	1.0E-06	NA	NA	NA	None	1	2.1E-16	1122	ESE	9.9E-12	736	W	1.4E-10	2
		- 2	planchetted, dry samples,	Pu-239	5.1E-09	1.0E-06		<u> </u>				5.1E-15							† -
			air filters and swipes	Pu-240	1.2E-09	1.0E-06						1.2E-15							+
			un meera una avripea	Pu-241	5.2E-08	1.0E-06						5.2E-14							+
				Pu-242	7.9E-14	1.0E-06						7.9E-20			1				+
			+	Am-241	2.5E-10	1.0E-06						2.5E-16							+
			+	U-238	2.8E-08	1.0E-06						2.8E-14	 						+
			+	U-235	1.3E-09	1.0E-06 1.0E-06						1.3E-15							+
				U-235	3.0E-08	1.0E-06 1.0E-06	-				-	3.0E-14							+
				U-234	3.UE-U8	1.0E-06	-				+	J.UE-14							+
252	17004	Dar · · A··	Crean alpha //	D., 220	2.15.10	1.05.00	NIA.	N.A.	N/A	Nie	1	2.15.10	1122	FCF	0.05.13	700	147	1 45 10	+ -
253	1708A	Room Air	Gross alpha/beta analysis of	Pu-238	2.1E-10	1.0E-06	NA	NA	NA	None	1	2.1E-16	1122	ESE	9.9E-12	736	W	1.4E-10	2
			planchetted, dry samples,	Pu-239	5.1E-09	1.0E-06						5.1E-15	-						
			air filters and swipes	Pu-240	1.2E-09	1.0E-06						1.2E-15							+
				Pu-241	5.2E-08	1.0E-06						5.2E-14							+
				Pu-242	7.9E-14	1.0E-06						7.9E-20							
				Am-241	2.5E-10	1.0E-06						2.5E-16							
				U-238	2.8E-08	1.0E-06						2.8E-14							
				U-235	1.3E-09	1.0E-06						1.3E-15							
				U-234	3.0E-08	1.0E-06						3.0E-14							

Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated	10 mrem/y 9	ite-Wide Dos	se Requirement	0.1 mrem/	y Monitoring	Requirement	Source
Dulluling	ROOM/ Area	Stack ID	Орегация	Radionaciaes	with Potential for	State	Height (m)	Diameter	Velocity	Device(s)	Abatement	Annual Emissions	Distance to		EDE	Distance	Direction	Unabated	Category
					Release (Ci)	Factor	3 4 ()	(m)	(m/s)	2 22(2)	Factor	(Ci)	SWMEI (m)	to SWMEI	(mrem)	to MEI (m)	to MEI	EDE (mrem)	1
253	1708B	Room Air	Gross alpha/beta analysis of	Pu-238	2.1E-10	1.0E-06	NA	NA	NA	None	1	2.1E-16	1122	ESE	9.9E-12	736	W	1.4E-10	2
			planchetted, dry samples,	Pu-239	5.1E-09	1.0E-06						5.1E-15							
			air filters and swipes	Pu-240	1.2E-09	1.0E-06						1.2E-15							
				Pu-241	5.2E-08	1.0E-06						5.2E-14							
				Pu-242	7.9E-14	1.0E-06						7.9E-20							
				Am-241 U-238	2.5E-10 2.8E-08	1.0E-06 1.0E-06						2.5E-16 2.8E-14							
				U-235	1.3E-09	1.0E-06						1.3E-15							
				U-234	3.0E-08	1.0E-06						3.0E-14							
				0 20 .	0.02 00							0.02							
253	1732	FHE-21	Flaming gross alpha/beta planchets	Pu-239	1.3E-13	1.0E+00	6.4	0.30	13.2	None	1	1.3E-13	1122	ESE	2.1E-11	736	W	1.4E-10	2
				Gross alpha	1.2E-13	1.0E+00						1.2E-13				798	WNW	1.4E-10	
				Gross beta	2.3E-13	1.0E+00						2.3E-13							
				H-3	1.1E-12	1.0E+00						1.1E-12							
0.50	1704		Bi dill di		0.75.10	1.05.00						0.75.40	1100	505	5.05.10	700	14/	7.05.44	
253	1734	Room Air	Distillation of environmental	H-3	6.7E-10	1.0E+00	NA	NA	NA	None	1	6.7E-10	1122	ESE	5.6E-12	736	W	7.8E-11	2
			samples	Gross alpha	5.4E-14 4.1E-13	1.0E+00 1.0E+00						5.4E-14 4.1E-13							
				Gross beta	4.1E-13	1.0E+00						4.1E-13							
253	1734	FGBE-1,2	Sieve soil samples	Gross alpha	2.7E-10	1.0E-06	6.1	0.10	23.6	HEPA	0.01	2.7E-18	1122	ESE	2.5E-16	736	W	2.4E-13	2
233	1754	1002 1,2	Sieve son samples	Gross beta	4.6E-10	1.0E-06	0.1	0.10	25.0	TILLY	0.01	4.6E-18	1122	LUL	2.32 10	730	**	2.42 13	
				5.555 beta															<u> </u>
253	1734	FHE-13	Samples and standards plating	Gross alpha	1.7E-11	1.0E+00	10.4	0.30	12.3	None	1	1.7E-11	1122	ESE	9.2E-11	798	WNW	4.6E-10	2
				Gross beta	2.2E-12	1.0E+00						2.2E-12							
				H-3	1.2E-11	1.0E+00						1.2E-11							
				Sr-90/Y-90	1.7E-12	1.0E+00						1.7E-12							
				Pu-239	7.8E-13	1.0E+00						7.8E-13							
253	1734	FHE-20	Quality control sample	Pu-239	2.5E-12	1.0E-03	10.4	0.30	12.3	None	1	2.5E-15	1122	ESE	1.8E-13	798	WNW	9.2E-13	2
			aliquoting	Sr/Y-90	2.2E-12	1.0E-03						2.2E-15							
				H-3	1.1E-10	1.0E-03						1.1E-13							
253	1734	FHE-11	Acid digestion for	H-3	6.8E-09	1.0E+00	10.4	0.30	12.3	None	1	6.8E-09	1122	ESE	2.8E-09	798	WNW	1.4E-08	2
233	1734	THE-TT	sample analysis	Gross alpha	3.4E-11	1.0E+00	10.4	0.50	12.3	None	'	3.4E-11	1122	LJL	2.01-03	736	W	1.4E-08	
			Sumple unarysis	Gross beta	2.2E-10	1.0E+00						2.2E-10				730		1.12.00	
				Sr/Y-90	2.8E-12	1.0E+00						2.8E-12							
				Pu-239	3.1E-12	1.0E+00						3.1E-12							
253	1910	FHE-22	Preparations of calibration	H-3	3.0E-11	1.0E-03	7.0	0.20	5.2	None	1	3.0E-14	1122	ESE	1.8E-15	736	W	2.1E-14	2
			standards	C-14	1.5E-11	1.0E-03						1.5E-14							
				P-32	1.5E-10	1.0E-03						1.5E-13							
D '11' OF		0 . 16 .1																	
			nducting bioassays and providing analyt		1 25 17	1.05.02	0.3	1.07	F 2	Nama	1	1 25 20	1020	FCF	2 75 17	1070	NINIE	1 45 16	-
254	108	FHE-1000	Analysis of urine for radionuclides	Am-243 Pu-242	1.3E-17 1.6E-16	1.0E-03 1.0E-03	8.2	1.07	5.3	None	1	1.3E-20 1.6E-19	1038	ESE	3.7E-17	1070 1055	NNE SW	1.4E-16 1.4E-16	2
			radiorideides	Pu-239	2.6E-17	1.0E-03						2.6E-20				849	WNW	1.4E-16	
				H-3	1.4E-14	1.0E-03						1.4E-17				013	WINV	1.46 10	
				Sr-90	3.1E-14	1.0E-03						3.1E-17							
				Y-90	3.1E-14	1.0E-03						3.1E-17							
254	109	FHE-1000	Analysis of urine for	Am-243	1.3E-17	1.0E-03	8.2	1.07	5.3	None	1	1.3E-20	1038	ESE	1.5E-18	1070	NNE	5.9E-18	2
			radionuclides										-						
254	110	FUE 1000	Amel size 5 d 5	A 2.11	0.05.10	1.05.00		1.07	F 2	NI	1	0.25.22	1000	FCE	1 25 11	1070	AINIE	F 0F 11	-
254	110	FHE-1000	Analysis of urine for radionuclides	Am-241 Am-243	8.2E-19 2.3E-17	1.0E-03 1.0E-03	8.2	1.07	5.3	None	1	8.2E-22 2.3E-20	1038	ESE	1.3E-11	1070	NNE	5.0E-11	2
			radionucildes			1.0E-03						2.3E-20 8.7E-21							
	1			('m_)///	Q 7E_10		1					1.1E-18	 						+
				Cm-244 Nn-237	8.7E-18 1 1F-15		1				1	1.12-10	I .	1 1		I .			1
				Np-237	1.1E-15	1.0E-03						8.9E-20							
				Np-237 Th-230	1.1E-15 8.9E-17	1.0E-03 1.0E-03						8.9E-20 8.0E-20							
				Np-237	1.1E-15	1.0E-03						8.9E-20 8.0E-20 2.7E-22							
				Np-237 Th-230 Cf-252	1.1E-15 8.9E-17 8.0E-17	1.0E-03 1.0E-03 1.0E-03						8.0E-20							
				Np-237 Th-230 Cf-252 U-233 U-234 U-235	1.1E-15 8.9E-17 8.0E-17 2.7E-19 2.6E-18 2.7E-19	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03						8.0E-20 2.7E-22 2.6E-21 2.7E-22							
				Np-237 Th-230 Cf-252 U-233 U-234 U-235 U-236	1.1E-15 8.9E-17 8.0E-17 2.7E-19 2.6E-18	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03						8.0E-20 2.7E-22 2.6E-21 2.7E-22 8.3E-20							
				Np-237 Th-230 Cf-252 U-233 U-234 U-235 U-236 U-238	1.1E-15 8.9E-17 8.0E-17 2.7E-19 2.6E-18 2.7E-19 8.3E-17 6.3E-21	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03						8.0E-20 2.7E-22 2.6E-21 2.7E-22 8.3E-20 6.3E-24							
				Np-237 Th-230 Cf-252 U-233 U-234 U-235 U-236 U-238 Mixed gamma	1.1E-15 8.9E-17 8.0E-17 2.7E-19 2.6E-18 2.7E-19 8.3E-17 6.3E-21 4.5E-12	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03						8.0E-20 2.7E-22 2.6E-21 2.7E-22 8.3E-20 6.3E-24 4.5E-15							
				Np-237 Th-230 Cf-252 U-233 U-234 U-235 U-236 U-238 Mixed gamma Cf-249	1.1E-15 8.9E-17 8.0E-17 2.7E-19 2.6E-18 2.7E-19 8.3E-17 6.3E-21 4.5E-12 6.7E-13	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03						8.0E-20 2.7E-22 2.6E-21 2.7E-22 8.3E-20 6.3E-24 4.5E-15 6.7E-16							
				Np-237 Th-230 Cf-252 U-233 U-234 U-235 U-236 U-238 Mixed gamma Cf-249 U-232	1.1E-15 8.9E-17 8.0E-17 2.7E-19 2.6E-18 2.7E-19 8.3E-17 6.3E-21 4.5E-12 6.7E-13 6.4E-13	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03						8.0E-20 2.7E-22 2.6E-21 2.7E-22 8.3E-20 6.3E-24 4.5E-15 6.7E-16 6.4E-16							
				Np-237 Th-230 Cf-252 U-233 U-234 U-235 U-236 U-238 Mixed gamma Cf-249 U-232 Po-209	1.1E-15 8.9E-17 8.0E-17 2.7E-19 2.6E-18 2.7E-19 8.3E-17 6.3E-21 4.5E-12 6.7E-13 6.4E-13 7.1E-14	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03						8.0E-20 2.7E-22 2.6E-21 2.7E-22 8.3E-20 6.3E-24 4.5E-15 6.7E-16 6.4E-16 7.1E-17							
				Np-237 Th-230 Cf-252 U-233 U-234 U-235 U-236 U-238 Mixed gamma Cf-249 U-232 Po-209 Pu-242	1.1E-15 8.9E-17 8.0E-17 2.7E-19 2.6E-18 2.7E-19 8.3E-17 6.3E-21 4.5E-12 6.7E-13 6.4E-13 7.1E-14 1.4E-13	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03						8.0E-20 2.7E-22 2.6E-21 2.7E-22 8.3E-20 6.3E-24 4.5E-15 6.7E-16 6.4E-16 7.1E-17							
				Np-237 Th-230 Cf-252 U-233 U-234 U-235 U-236 U-238 Mixed gamma Cf-249 U-232 Po-209 Pu-242 Pu-239	1.1E-15 8.9E-17 8.0E-17 2.7E-19 2.6E-18 2.7E-19 8.3E-17 6.3E-21 4.5E-12 6.7E-13 6.4E-13 7.1E-14 1.4E-13 2.2E-14	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03						8.0E-20 2.7E-22 2.6E-21 2.7E-22 8.3E-20 6.3E-24 4.5E-15 6.7E-16 6.4E-16 7.1E-17 1.4E-16 2.2E-17							
				Np-237 Th-230 Cf-252 U-233 U-234 U-235 U-236 U-238 Mixed gamma Cf-249 U-232 Po-209 Pu-242 Pu-239 P-32	1.1E-15 8.9E-17 8.0E-17 2.7E-19 2.6E-18 2.7E-19 8.3E-17 6.3E-21 4.5E-12 6.7E-13 6.4E-13 7.1E-14 1.4E-13 2.2E-14 7.0E-12	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03						8.0E-20 2.7E-22 2.6E-21 2.7E-22 8.3E-20 6.3E-24 4.5E-15 6.7E-16 6.4E-16 7.1E-17 1.4E-16 2.2E-17 7.0E-15							
				Np-237 Th-230 Cf-252 U-233 U-234 U-235 U-236 U-238 Mixed gamma Cf-249 U-232 Po-209 Pu-242 Pu-239	1.1E-15 8.9E-17 8.0E-17 2.7E-19 2.6E-18 2.7E-19 8.3E-17 6.3E-21 4.5E-12 6.7E-13 6.4E-13 7.1E-14 1.4E-13 2.2E-14	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03						8.0E-20 2.7E-22 2.6E-21 2.7E-22 8.3E-20 6.3E-24 4.5E-15 6.7E-16 6.4E-16 7.1E-17 1.4E-16 2.2E-17							

Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated	10 mrem/y 5	Site-Wide Do	se Requirement	0.1 mrem	y Monitoring	Requirement	Source
					with Potential for	State	Height (m)	Diameter	Velocity	Device(s)	Abatement	Annual Emissions	Distance to		EDE	Distance	Direction	Unabated	Category
254	110	(1)		1.125	Release (Ci)	Factor		(m)	(m/s)		Factor	(Ci)	SWMEI (m)	to SWMEI	(mrem)	to MEI (m)	to MEI	EDE (mrem)	
254	110	(continued)		I-125	9.0E-13	1.0E-03						9.0E-16							
				Sr-90	5.0E-14	1.0E-03						5.0E-17							
				Y-90	5.0E-14	1.0E-03						5.0E-17							
				Np-237	1.1E-10 6.2E-10	1.0E-03 1.0E-03						1.1E-13 6.2E-13		-					
				Np-239	9.1E-13	1.0E-03						9.1E-16							
				Cm-242 Th-230	9.1E-13 9.4E-12	1.0E-03						9.4E-15							
				Cf-252	9.4E-12 8.4E-12	1.0E-03						9.4E-15 8.4E-15							
				U-233	2.2E-16	1.0E-03						2.2E-19							
				U-234	8.2E-15	1.0E-03						8.2E-18							
				U-235	2.3E-16	1.0E-03						2.3E-19							
				U-236	5.8E-14	1.0E-03						5.8E-17							
				U-238	2.0E-15	1.0E-03						2.0E-18							
				0 230	2.02 13	1.02 03						2.02.10							
254	113	FHE-1000	Analysis of urine for	Pu-242	1.6E-16	1.0E-03	8.2	1.07	5.3	None	1	1.6E-19	1038	ESE	3.8E-17	1070	NNE	1.4E-16	2
231	113	1112 1000	radionuclides	Pu-239	2.6E-17	1.0E-03	0.2	1.07	5.5	Hone		2.6E-20	1030	LOL	3.0L 17	1055	SW	1.4E-16	
			- Tudioridones	H-3	1.4E-14	1.0E-03						1.4E-17				817	W	1.4E-16	
				C-14	1.4E-14	1.0E-03						1.4E-17				849	WNW	1.4E-16	
				Sr-90	3.1E-14	1.0E-03						3.1E-17				0.0			
				Y-90	3.1E-14	1.0E-03						3.1E-17							
				Am-243	1.3E-17	1.0E-03						1.3E-20							
				7.111 2.10		1102 00													
Building 255	is operated by Haz	ards Control and houses a ra	diation calibration and standards labora	tory. Many operation	ns involve the use of	sealed sources	5.												
255	165	FHE-4	Analysis of urine for	I-125	2.3E-09	1.0E-03	6.9	0.30	5.1	None	1	2.3E-12	1056	E	6.1E-12	790	W	1.8E-11	2
255	165	FHE-4					6.9	0.30	5.1	None	I		1056	E	6.1E-12	790	VV	1.8E-11	
			radionuclides	I-131 Th-230	7.2E-09 5.7E-14	1.0E-03						7.2E-12 5.7E-17		-					
						1.0E-03													
				Th-232	1.0E-16	1.0E-03						1.0E-19							
				U-233	1.0E-11 1.3E-15	1.0E-03 1.0E-03						1.0E-14 1.3E-18		-					
				U-238 Np-237	5.7E-14	1.0E-03						5.7E-17							
				Cm-244	3.8E-14 3.8E-15	1.0E-03						3.8E-17 3.8E-18							
				Am-241		1.0E-03						1.9E-17		-					
				Am-243	1.9E-14 1.9E-14	1.0E-03						1.9E-17 1.9E-17							
				Pu-239 Pu-242	1.9E-14 1.9E-15	1.0E-03 1.0E-03						1.9E-17 1.9E-18							
				Fu-242	1.95-13	1.02-03						1.95-16							
255	180	FHE-2	Tritium gas monitor calibrations	H-3	2.5E-02	1.0E+00	8.1	0.31	5.2	None	1	2.5E-02	1056	E	9.9E-06	790	W	3.9E-05	2
Ruilding 281	is part of the Energ	ay and Environment Director	ate. Tracer work, dissolution studies an	d flow studies are co	nducted in this buildir	20													
building 201	is part of the Lifery	gy and Environment Directors	ate. Tracer work, dissolution studies an	la now studies are co	Triducted III triis buildii	ig.													
281	1174	FHE-13	Tracer work	Ni-63	1.0E-05	1.0E-03	6.7	0.30	6.1	None	1	1.0E-08	1332	ESE	2.4E-11	579	NNE	3.1E-10	2
281	1305	Room air	Dissolution studies	U-238	4.3E-09	1.0E-03	NA	NA	NA	None	1	4.3E-12	1332	ESE	1.0E-10	753	WNW	1.8E-09	2
281	1307	FHE-6	Tracer work	Np-237	2.5E-12	1.0E-03	6.4	0.61	2.7	None	1	2.5E-15	1332	ESE	4.0E-08	753	WNW	5.6E-07	2
201	1307	FHE-0	Tracer work	U-238	4.10E-14	1.0E-03	0.4	0.61	2.1	None	· ·	4.1E-17	1332	ESE	4.0E-08	733	VVINVV	3.0E-07	
				U-235	5.28E-16	1.0E-03						5.3E-19		+					
				U-234	3.83E-15	1.0E-03						3.8E-18							
				Pu-239	3.3E-07	1.0E-03						3.3E-10							
				Pu-242	2.1E-11	1.0E-03						2.1E-14							
				U-233	2.3E-08	1.0E-03						2.3E-11							
				Pu-244	1.6E-09	1.0E-03						1.6E-12							
				Ni-63	2.0E-04	1.0E-03						2.0E-07							
				Ni-59	7.0E-08	1.0E-03						7.0E-11							
				Tc-99	1.0E-07	1.0E-03						1.0E-10							
				Sr-90	1.0E-05	1.0E-03						1.0E-08							
				Ca-41	1.0E-04	1.0E-03						1.0E-07							
				Be-10	1.0E-05	1.0E-03						1.0E-08							
				Pu-239/U-233	1.0E-07	1.0E-03						1.0E-10							
201	1211	FUE 10	Colorina	0.14	1.05.04	1.05.00	C 1	0.41	4.0	Man -	1	1.05.07	1222	FCF	4.05.00	750	\A/A I\A/	E	1
281	1311	FHE-12	Solution preparation	C-14	1.9E-04	1.0E-03	6.1	0.41	4.0	None	1	1.9E-07	1332	ESE	4.0E-09	753	WNW	5.5E-08	2
			+	Cl-36 H-3	1.0E-05 2.5E-05	1.0E-03 1.0E-03						1.0E-08 2.5E-08							+
				11-3	2.31-03	1.01-03						2.31-00							
					8.0E-08	1.0E-03	6.7	0.30	6.1	None	1	8.0E-11	1332	ESE	6.2E-09	579	NNE	8.3E-08	2
281	1323	FHE-1	Radioactivity migration studies	Na-22	0.01 00						1	1.2E-10				T			
281	1323	FHE-1	Radioactivity migration studies	Na-22 U-238	1.2E-07	1.0E-03						1.25-10							
281	1323	FHE-1	Radioactivity migration studies			1.0E-03 1.0E-03						5.5E-12							
281	1323	FHE-1	Radioactivity migration studies	U-238	1.2E-07														
				U-238 U-235 U-234	1.2E-07 5.5E-09 1.3E-07	1.0E-03						5.5E-12							
			Radioactivity migration studies torate. Residual contamination exists in	U-238 U-235 U-234	1.2E-07 5.5E-09 1.3E-07	1.0E-03						5.5E-12							
				U-238 U-235 U-234	1.2E-07 5.5E-09 1.3E-07	1.0E-03	NA	NA	NA	None	1	5.5E-12	1332	ESE	6.2E-13	753	WNW	1.1E-11	2

Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated	<u>10 mrem/y</u> :	Site-Wide Do	se Requirement	<u>0.1 mrem/</u>	y Monitoring	Requirement	Source
					with Potential for	State	Height (m)	Diameter	Velocity	Device(s)	Abatement	Annual Emissions	Distance to	Direction	EDE	Distance	Direction	Unabated	Categor
					Release (Ci)	Factor	<u> </u>	(m)	(m/s)		Factor	(Ci)	SWMEI (m)	to SWMEI	(mrem)	to MEI (m)	to MEI	EDE (mrem)	
Building 29	2 is administered by	the Environmental Programs D	irectorate. Residual contamination exi	ists throughout the f	acility from the past	operation of a	a rotating target i	neutron source.											
292	1200,1202	Room Air	Contamination	H-3	1.8E+00	1.0E-03	NA	NA	NA	None	1	1.8E-03	1380	ESE	3.9E-06	655	W	9.2E-05	2
LJL	1204	Room Air	Contamination	H-3	2.3E+01	1.0E-03	NA NA	NA NA	NA NA	None	1	2.3E-02	1300	LJL	3.3L-00	033	VV	3.2L-03	+
	1402, 1402A	Room Air	Contamination	H-3	1.8E+00	1.0E-03	NA NA	NA NA	NA	None	1	1.8E-03							1
	1404, 1406																		
	1407																		
						L													
Building 29	8 is part of the Lase	er Fusion Program. Small amoui	nts of tritium are used in this facility in	n conjunction with fus	sion target research a	and developm	ent.												
298	160	Room Air	D-T layering experiment	H-3	4.0E-03	1.0E+00	NA	NA	NA	None	1	4.0E-03	1398	SE	5.7E-07	264	NNE	3.0E-05	2
230	100	Noom All	D 1 layering experiment	113	4.0L 03	1.02+00	IVA	INA	ING	None		4.02.03	1550	JL.	5.7 L 07	201	ININE	3.02.03	+
298	189	FHE-14	Laser fusion target coating	U-238	1.3E-04	1.0E-03	6.4	0.63	15.1	HEPA	0.01	1.3E-09	1398	SE	1.9E-08	344	NE	6.3E-05	2
298	Various	Room Air	Laser fusion target research	H-3	1.0E-03	1.0E+00	NA	NA	NA	None	1	1.0E-03	1398	SE	1.4E-07	264	NNE	7.6E-06	2
			and development																
Puildings 2	21 221 A 221 D and	d 221C are the Material Eabrica	tion Shops and are part of the Machar	nical Engineering Don	artmont Operations	in this comple	ov include milling	chaning and ma	hining of doplot	od uranjum									+
			tion Shops and are part of the Mechar be moved from machine to machine.								nends								+
			NOTE: Machining only occurs in 321C.		dramam parts occasi	dially dilucing	Theat treatment	. The amount of	depicted drama	III triat is rialidica de	perius								+
			PA/DOE Memorandum of Understandin																1
		•																	
321A	1001A	FHE-24	Machining and manufacturing	U-234	7.5E-04	1.0E-06	3.7	0.46	2.9	HEPA	0.01	7.5E-12	1032	ENE	1.10E-08	326	SW	8.3E-06	2
				U-235	1.0E-04	1.0E-06						1.0E-12							
				U-238	8.1E-03	1.0E-06	-					8.1E-11							+
321C	234B	FHE-13	Lapping of DU metal	U-238	1.6E-04	1.0E-06	10.7	0.49	2.5	None	1	1.6E-10	1032	ENE	1.80E-08	326	SW	4.2E-08	2
3210	2346	FRE-13	Lapping of Do metal	U-235	2.0E-06	1.0E-06	10.7	0.49	2.5	None	'	2.0E-12	1032	EINE	1.00E-00	326	SVV	4.26-00	+
				U-234	9.4E-06	1.0E-06						9.4E-12							+
				0 20 .	0.12.00	1.02.00						0112 12							1
321C	Various**	FHE-9	Machining and manufacturing	U-234	3.2E+00	1.0E-06	8.5	0.31	16.1	HEPA	0.01	3.2E-08	1032	ENE	3.4E-08	252	SW	6.2E-06	2
		FHE-11		U-235	4.0E-02	1.0E-06	12.5	0.60	6.0	HEPA	0.01	4.0E-10							
		FHE-15		U-238	3.0E-01	1.0E-06	11.2	0.23	13.4	HEPA	0.01	3.0E-09							
		FEV-1000					11.3	0.83	6.5	HEPA	0.01								
Duilding 21	2 is approted by the	Machaniaal Engineering Danast	tment			-													+
bulluling 32	.z is operated by the	Mechanical Engineering Depart	inent.																+
322	109	FHE-1	Cleaning and plating	U-234	3.1E-07	1.0E-06	7.9	0.35	1.0	None	1	3.1E-13	930	ENE	5.0E-10	416	SW	1.8E-09	2
022			of depleted uranium	U-235	4.3E-08	1.0E-06	1.10	0.00			·	4.3E-14	- 555		0.02 10		0	1.02 00	+-
			·	U-238	3.3E-06	1.0E-06						3.3E-12							
Building 32	7 is operated by the	Mechanical Engineering Depart	tment.																
227	1275	Danier Air	Non destructive observation	U-234	1.3E-05	1.0E-06	NA	NA	NIA	Ness	1	1.3E-11	1018	ENE	1.9E-08	425	SW	1.2E-07	2
327	1275	Room Air	Non-destructive ultrasonic material evaluation	U-235	1.9E-06	1.0E-06	INA	INA	NA	None	'	1.9E-12	1016	EINE	1.9E-06	423	SVV	1.26-07	+
			material evaluation	U-238	1.4E-04	1.0E-06						1.4E-10							+
																			1
Building 33	1 is operated by the	Defense and Nuclear Technolo	gies Directorate. The building houses	the tritium research	facility and associate	d laboratories	S.												
			continuously monitored in compliance		ations. Monitoring da	ta, rather tha	n the inventory a	pproach, are use	d to determine e	emissions.									
			PA/DOE Memorandum of Understanding emissions as HTO, as directed by U.S		door from UT and U	ITO amissions	anleylated appro	priotoly using the	NEW/TRIT mode	Lie E CE O2 Coe dies	augaian in Castian \	/III. subsection "Med	laling Doos from	a Tritiuma "					+
Calcula	ed dose of 6.1E-03	mrem includes modeling the Hi	emissions as ATO, as directed by 0.5	EPA, Region IA. The	dose irom ni and n		calculated appro	priately using the	NEWIKII Mode	1 IS 3.0E-US. See uist	Lussion in Section V	iii, subsection Mou	leling bose from	i iridum.					+
331	All**	Stack 1	Tritium research and development	H-3	*	1.0E+00	30.0	1.22	7.6	None	1	2.5E-02	957	ENE	8.1E-03	957	ENE	8.1E-03	3
		Stack 2	Decontamination of parts	H-3	*	1.0E+00	30.0	1.22	10.5	None	1	3.6E+01			***5.6E-03			***5.6E-03	
			plutonium research. Exhausts from gl			L													
	, , , ,	-,	rs. Exhausts are monitored with both		1. 3 (.,		-,		(/	The air menitarian	ata for all arries:	nointe							+
			ciated with specific tasks is classified, w the limit of sensitivity of the analyti		rs approach, based o	ni inventory, (Carinot de Utilizeo	without classify	ing this report.	rne all monitoring da	ata for all emission	μυπιτο							+
			bated EDE cannot be determined (see		n II, subsection "Stack	ı k Monitorina f	or Gross Alnha ar	nd Gross Beta Rad	diation.")										+
	, g piu		(000		,				,										1
332	Increment 1	FHE-1000/2000	Plutonium research	Transuranics	*	NA	8.8	0.8x1.1	17.3	Double HEPA	0.000001	0.0E+00	912	ENE	0.0E+00	**	**	**	3
	Rooms																		
222	la and the state of	FODE 1000/0000	District	Tuesta	<u>.</u>	A.I.A.	11	0.2		T2-1- U554	0.000001	0.05.00	010	ENE	0.05.00	**	**	11	+
332	Increment 1 Glove boxes	FGBE-1000/2000	Plutonium research	Transuranics	*	NA	11	0.3	6.9	Triple HEPA	0.000001	0.0E+00	912	ENE	0.0E+00	**	**	**	3
	Giove boxes																		+
332	Downdraft	FHE-4/5	Plutonium research	Transuranics	*	NA	11	0.2	14.2	Double HEPA	0.0001	0.0E+00	912	ENE	0.0E+00	**	**	**	3
552	Sommatt	.112 1/3	. iscomani rescaren	ariour arinos		19/1	† ··	J.L		DOUBLE TILL A	0.0001	5.02100	312	E11E	3.52.100				+ -
332	Loft	FE-4	Plutonium research	Transuranics	*	NA	11	0.6x0.9	4.6	HEPA	0.01	0.0E+00	912	ENE	0.0E+00	**	**	**	3
JJL		FE-5	Plutonium research	Transuranics	*	NA	11	0.6x0.9	4.6	HEPA	0.01	0.0E+00	912	ENE	0.0E+00	**	**	**	3
332																			
								-						_					
332	Increment 1	FGBE-3000/4000	Plutonium research	Transuranics	*	NA	11	0.3	2	Triple HEPA	0.000001	0.0E+00	912	ENE	0.0E+00	**	**	**	3
	Increment 1 Glove boxes	FGBE-3000/4000	Plutonium research	Transuranics	*	NA	11	0.3	2	Triple HEPA	0.000001	0.0E+00	912	ENE	0.0E+00	**	**	**	3

Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated	10 mrem/y S	ite-Wide Dos	se Requirement	0.1 mrem/	y Monitoring	Requirement	Source
					with Potential for	State	Height (m)	Diameter	Velocity	Device(s)	Abatement	Annual Emissions	Distance to	Direction	EDE	Distance	Direction	Unabated	Category
332	Increment 3	FFE-1000/2000	Plutonium research	Transuranics	Release (Ci)	Factor NA	10.1	(m) 0.9	(m/s) 12.2	Room—Double HEPA	Factor 0.000001	(Ci) 0.0E+00	SWMEI (m) 912	to SWMEI ENE	(mrem) 0.0E+00	to MEI (m)	to MEI	EDE (mrem) **	3
552	Room and	FGBE-7000/8000	. iacomanii roodardii			.,,,		0.0		Glove Box—Triple HEP		5.52100	312	2.32	0.02100				
	Glove boxes																		
Building 34	11 is a Lasers Direct	orate facility.																	
341	1107	Room Air	Blower decontamination	U-238	9.6E-10	1.00E-03	NA	NA	NA	None	1	9.6E-13	872	E	1.1E-10	770	SW	2.90E-10	2
				U-235 U-234	1.2E-11 8.9E-11	1.00E-03						1.2E-14 8.9E-14				591	SSW	2.90E-10	
				0-234	0.9E-11	1.00E-03						0.9E-14							+
			arch Directorate includes Buildings 361																
			mal research, and incorporated in anim exhausted. Most of the organs that co																
			n-14, phosphorous-32, phosphorous-3						ids.										
361	1020	Room Air	DNA hybridization	P-32	1.3E-03	1.0E-03	NA	NA	NA	None	1	1.3E-06	918	ESE	2.1E-08	976	W	1.1E-07	2
361	1137	Room Air	Protein hybridization	P-32	4.7E-02	1.0E-03	NA	NA	NA	None	1	4.7E-05	918	ESE	9.1E-07	976	W	4.6E-06	1
361	1238	Room Air	P-32 Labeling	P-32	8.0E-05	1.0E-03	NA	NA	NA	None	1	8.0E-08	918	ESE	1.3E-09	976	W	7.0E-09	2
301	1236	KOOIII AII	r-32 Labelling	F-32	8.0E-03	1.0E-03	NA NA	INA	INA	None		8.0E-08	310	ESE	1.3E-03	370	VV	7.0E-09	
361	1445	Room Air	Radiolabeling of DNA substrates	P-32 S-35	2.2E-04 4.1E-04	1.0E-03 1.0E-03	NA	NA	NA	None	1	2.2E-07 4.1E-07	918	ESE	6.2E-09	976	W	3.4E-08	2
361	1446	FHE-15	Radiolabeling of DNA substrates	P-32	5.5E-04	1.0E-03	6.2	0.42	1.7	None	1	5.5E-07	918	ESE	8.7E-09	976	W	4.7E-08	2
361	1542	FHE-12	Hybridization and enzyme assay	P-32	2.0E-07	1.0E-03	7.0	0.41	4.4	None	1	2.0E-10	918	ESE	2.9E-12	976	W	1.5E-11	2
361	1546	FHE-10	DNA protein interaction studies	P-32	3.2E-04	1.0E-03	1.7	0.41	0.5	None	1	3.2E-07	918	ESE	4.9E-09	976	W	2.5E-08	2
361	1664	Room Air	DNA protein interaction studies DNA hybridization	P-32	6.4E-04	1.0E-03	NA	NA	NA	None	1	6.4E-07	918	ESE	1.0E-08	976	W	5.6E-08	2
361	1664	KOOM AII	DINA hybridization	P-32	6.4E-04	1.0E-03	NA	NA NA	NA NA	None	I	6.4E-07	918	ESE	1.0E-08	976	VV	5.6E-U8	
361	1664A	Room Air	Labeling Olegios for high density filter hybridization	P-32	4.3E-04	1.0E-03	NA	NA	NA	None	1	4.3E-07	918	ESE	8.4E-09	976	W	4.2E-08	1
0.01	1740	EUE O		5.00	0.05.04	1.05.00	7.0	0.44				0.05.07	010	505	2.25.22	070	144	1.75.00	
361	1742	FHE-8	DNA hybridization	P-32	2.2E-04	1.0E-03	7.0	0.41	4.4	None	1	2.2E-07	918	ESE	3.3E-09	976	W	1.7E-08	2
361	1846	Room Air	Human genome research	P-32	2.6E-04	1.0E-03	NA NA	NA NA	NA	None	1	2.6E-07	918	ESE	4.2E-09	976	W	2.3E-08	2
Building 36		EUE 1000	Comment of Station In URI C	11.2	1.05.04	1.05.02	6.0	0.65	2.7	News		1.05.07	000	FOE	1 45 00	002	14/	0.45.00	
362	105	FHE-1000	Compound purification by HPLC	H-3 C-14	1.0E-04 1.0E-04	1.0E-03 1.0E-03	6.8	0.65	2.7	None	1	1.0E-07 1.0E-07	992	ESE	1.4E-09	893	W	9.4E-09	2
362	106	FHE-1000	Characterization of	C-14	1.0E-08	1.0E-03	6.8	0.65	2.7	None	1	1.0E-11	992	ESE	6.7E-09	893	W	4.6E-08	2
302	100	FRE-1000	metabolic pathways	H-3	5.0E-04	1.0E-03	0.0	0.03	2.1	None		5.0E-07	332	ESE	0.7 E=03	093	VV	4.02-00	
Duilding 20	22																		1
Building 36 363	1009	FHE-2000	Human urine sample project	H-3	1.0E-09	1.0E-03	1.7	0.41	0.4	HEPA	0.01	1.0E-14	1000	ESE	1.5E-16	888	W	1.2E-13	2
				C-14	1.0E-09	1.0E-03						1.0E-14							
363	1010	Room Air	HPLC analysis	H-3	1.0E-09	1.0E-03	NA	NA	NA	None	1	1.0E-12	1000	ESE	1.6E-14	888	W	1.3E-13	2
				C-14	1.0E-09	1.0E-03						1.0E-12							
Building 36																			
364	1509	FHE-02P	AMS sample preparation	H-3	5.5E-14	1.0E+00	5.5	0.52	2.9	None	1	5.5E-14	987	ESE	8.6E-09	912	W	6.8E-08	2
				C-14	5.5E-07	1.0E+00						5.5E-07							
364	1509A	Room Air	AMS sample preparation	H-3 C-14	5.5E-14 5.5E-07	1.0E+00 1.0E+00	NA	NA	NA	None	1	5.5E-14 5.5E-07	987	ESE	7.9E-09	912	W	5.6E-08	2
		_																	1
364	1519	Room Air	DNA and protein extraction	C-14 H-3	5.0E-06 5.0E-06	1.0E-03 1.0E-03	NA	NA	NA	None	1	5.0E-09 5.0E-09	987	ESE	8.0E-11	912	W	6.3E-10	2
Building 36	35																		<u> </u>
365	104	FHE-1000	Equipment decontamination	C-14	1.0E-09	1.0E-03	6.1	0.58	7.2	HEPA	0.01	1.0E-14	991	ESE	1.2E-16	902	W	6.1E-14	2
				H-3	1.0E-09	1.0E-03						1.0E-14							
365	109	FHE-5	Animal housing	C-14	1.3E-05	1.0E-03	1.7	0.41	0.6	Double HEPA	0.0001	1.3E-12	991	ESE	2.0E-14	902	W	1.6E-09	2
		2 0	a riodoniy	H-3	5.0E-08	1.0E-03		J1	3.3		3.0001	5.0E-15	331		11				
Building 36	66																		+
366	111	Room Air	Labeling	P-32	2.0E-03	1.0E-03	NA	NA	NA	None	1	2.0E-06	925	ESE	3.2E-08	998	W	1.7E-07	2

Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated		Site-Wide Dos				Requirement	Source
					with Potential for Release (Ci)	State Factor	Height (m)	Diameter (m)	Velocity (m/s)	Device(s)	Abatement Factor	Annual Emissions (Ci)	Distance to SWMEI (m)		(mrem)	Distance to MEI (m)	Direction to MEI	Unabated EDE (mrem)	Categor
uildina 3	378 is part of the En	ergy and Environment Directorate	 e. Small quantities of radioactive trace	ı ers are handled in thi	. ,	ractor	+	(111)	(111/5)		ractor	(CI)	SWIVIEI (III)	LO SVVINEI	(miem)	LO MEI (III)	LO IVIEI	EDE (IIIrem)	1
378	105	FHE-1,11	Tracer work	Am-243	9.2E-12	1.0E-03	8.5	0.30	5.8	None	1	9.2E-15	875	ESE	4.2E-12	1041	W	1.9E-11	2
				Pu-239	1.5E-14	1.0E-03						1.5E-17							
				Pu-242	1.5E-12	1.0E-03						1.5E-15							
				U-233	4.6E-11	1.0E-03						4.6E-14							
378	120	FHE-1,2,3,4,5,8,9,10	Tracer work	Am-241	1.2E-08	1.0E-03	8.5	0.30	5.9	None	1	1.2E-11	875	ESE	2.6E-09	1041	W	1.2E-08	2
310	120	1112 1,2,3,1,3,5,3,10	Tracer work	Am-243	7.7E-11	1.0E-03	0.5	0.50	3.3	Hone		7.7E-14	0.3	LOL	2.02 03	1011	•	1.22 00	-
				Cd-109	7.7E-09	1.0E-03						7.7E-12							
				Co-57	3.1E-10	1.0E-03						3.1E-13							
				Co-60	1.2E-08	1.0E-03						1.2E-11							
				Cs-134	1.5E-07	1.0E-03						1.5E-10							
				Cs-137	1.5E-08	1.0E-03 1.0E-03						1.5E-11 3.1E-16							_
				Np-237 Pu-239	3.1E-13 1.5E-11	1.0E-03						1.5E-14							+
				Pu-240	1.2E-11	1.0E-03						1.2E-14							
				Pu-242	1.5E-12	1.0E-03						1.5E-15							
				Pu-244	7.7E-11	1.0E-03						7.7E-14							
				Sr-85	1.4E-08	1.0E-03						1.4E-11							
				U-238	1.5E-13	1.0E-03						1.5E-16							
				U-235	6.8E-15	1.0E-03	-					6.8E-18							+
	+			U-234 U-238	1.6E-13 2.8E-13	1.0E-03 1.0E-03	-					1.6E-16 2.8E-16							+
				U-235	3.6E-15	1.0E-03						3.6E-18							+
				U-234	2.6E-14	1.0E-03						2.6E-17							
ıilding 4	191 was part of the I	Uranium Atomic Vapor Laser Isoto	ope Separation (U-AVLIS) program, op	erated by The United	d States Enrichment	Corporation (JSEC). In June 19	99, USEC susper	nded further dev	elopment of the U-AV	LIS technology.								
			n-series high efficiency particulate (HE			L	L	==											
			r atmospheric discharge points, althou							ns.									
Becaus	e monitoring takes p	liace after HEPA flitration, an unai	pated EDE cannot be determined (see	discussion in Section	n II, subsection "Stac	K Monitoring 1	or Gross Alpha ar	id Gross Beta Kai	diation.")										
			0	0	*	NA	9.1	0.9	12.1	Double HEPA	0.0001	0.0E+00	1000	SSE	0.0E+00	**	**	**	3
491	All	l FFE-1	Uut of Service	uross albha															_
491	All	FFE-1	Out of service	Gross alpha Gross beta	*	IVA		0.5	12.1	Bouble HELFY	3.333	0.0E+00							
-				Gross beta	*														
uilding 5	513 is operated by the	he Radioactive and Hazardous Wa	ste Management Division. The Stabili:	Gross beta zation Unit is a mech	* nanized mixing device	used to make	e homogeneous m	nixtures of waste	e. Solidification a	gents are added duri	ng mixing to transf	er sludges to solids							
ilding 5	513 is operated by the	he Radioactive and Hazardous Wa		Gross beta zation Unit is a mech	* nanized mixing device	used to make	e homogeneous m	nixtures of waste	e. Solidification a	gents are added duri	ng mixing to transf	er sludges to solids							
ilding 5	513 is operated by the	he Radioactive and Hazardous Wa	ste Management Division. The Stabili:	Gross beta zation Unit is a mech	* nanized mixing device	used to make	e homogeneous m	nixtures of waste	e. Solidification a	gents are added duri	ng mixing to transf	er sludges to solids							
ilding S	513 is operated by the	he Radioactive and Hazardous Wa	ste Management Division. The Stabili:	Gross beta zation Unit is a mech	* nanized mixing device	used to make	e homogeneous m	nixtures of waste	e. Solidification a	gents are added duri	ng mixing to transf	er sludges to solids	528	NE	3.4E-07	217	SW	8.3E-07	1
ilding 5 e Micro	513 is operated by the offiltration Unit filters	ne Radioactive and Hazardous Wa out waste radioactive particles.	ste Management Division. The Stabili In the Laboratory, small quantities of	Gross beta zation Unit is a mech waste materials are s	* lanized mixing device sampled, treated, and	used to maked stored. No	e homogeneous m releases are assu	ixtures of waste med to occur fro	e. Solidification a om waste storag	gents are added duri e because the wastes	ng mixing to transl are fully containe	er sludges to solids d.		NE	3.4E-07	217	SW	8.3E-07	1
ilding S	513 is operated by the offiltration Unit filters	ne Radioactive and Hazardous Wa out waste radioactive particles.	ste Management Division. The Stabili In the Laboratory, small quantities of Treatment of hazardous, mixed	Gross beta zation Unit is a mech waste materials are s	* lanized mixing device sampled, treated, and 4.1E-04	used to maked stored. No	e homogeneous m releases are assu	ixtures of waste med to occur fro	e. Solidification a om waste storag	gents are added duri e because the wastes	ng mixing to transl are fully containe	er sludges to solids d. 4.1E-07		NE	3.4E-07	217	SW	8.3E-07	1
ilding S	513 is operated by the offiltration Unit filters	ne Radioactive and Hazardous Wa out waste radioactive particles.	ste Management Division. The Stabili In the Laboratory, small quantities of Treatment of hazardous, mixed	Gross beta zation Unit is a mech waste materials are s H-3 U-234	* anized mixing device sampled, treated, and 4.1E-04 5.1E-08	used to make distored. No 1.0E-03 1.0E-03	e homogeneous m releases are assu	ixtures of waste med to occur fro	e. Solidification a om waste storag	gents are added duri e because the wastes	ng mixing to transl are fully containe	er sludges to solids d. 4.1E-07 5.1E-11		NE	3.4E-07	217	SW	8.3E-07	1
ne Micro	513 is operated by the filtration Unit filters Stabilization	he Radioactive and Hazardous Wa out waste radioactive particles. Room Air	ste Management Division. The Stabilis In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste	Gross beta zation Unit is a mech waste materials are s H-3 U-234 U-235 U-238	* nanized mixing device sampled, treated, and 4.1E-04 5.1E-08 7.1E-09 5.5E-07	used to maked stored. No 1.0E-03 1.0E-03 1.0E-03 1.0E-03	e homogeneous m releases are assu NA	ixtures of waste med to occur fro NA	e. Solidification a om waste storag	gents are added duri e because the wastes None	ng mixing to transls are fully containe	4.1E-07 5.1E-11 7.1E-12 5.5E-10	528						
ne Micro	513 is operated by the offiltration Unit filters	ne Radioactive and Hazardous Wa out waste radioactive particles.	ste Management Division. The Stabilis In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste	Gross beta zation Unit is a mech waste materials are s H-3 U-234 U-235 U-238	* lanized mixing device sampled, treated, and 4.1E-04 5.1E-08 7.1E-09 5.5E-07	1.0E-03 1.0E-03 1.0E-03	e homogeneous m releases are assu	ixtures of waste med to occur fro	e. Solidification a om waste storag	gents are added duri e because the wastes	ng mixing to transl are fully containe	4.1E-07 5.1E-11 7.1E-12 5.5E-10		NE NE	3.4E-07 8.7E-09	217	SW	8.3E-07 1.3E-08	
ne Micro	513 is operated by the filtration Unit filters Stabilization	he Radioactive and Hazardous Wa out waste radioactive particles. Room Air	ste Management Division. The Stabilis In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste	Gross beta zation Unit is a mech waste materials are s H-3 U-234 U-235 U-238 I-125 I-131	* lanized mixing device sampled, treated, and 4.1E-04 5.1E-08 7.1E-09 5.5E-07 7.8E-07 2.8E-08	used to maked a stored. No 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	e homogeneous m releases are assu NA	ixtures of waste med to occur fro NA	e. Solidification a om waste storag	gents are added duri e because the wastes None	ng mixing to transls are fully containe	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13	528						
ne Micro	513 is operated by the filtration Unit filters Stabilization	he Radioactive and Hazardous Wa out waste radioactive particles. Room Air	ste Management Division. The Stabilis In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste	Gross beta zation Unit is a mech waste materials are s H-3 U-234 U-235 U-238 I-125 I-131 Cs-137	* anized mixing device sampled, treated, and 4.1E-04 5.1E-08 7.1E-09 5.5E-07 7.8E-07 2.8E-08 2.5E-07	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	e homogeneous m releases are assu NA	ixtures of waste med to occur fro NA	e. Solidification a om waste storag	gents are added duri e because the wastes None	ng mixing to transls are fully containe	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13 2.5E-12	528						
illding 5 ie Micro	513 is operated by the filtration Unit filters Stabilization	he Radioactive and Hazardous Wa out waste radioactive particles. Room Air	ste Management Division. The Stabilis In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste	Gross beta zation Unit is a mech waste materials are s H-3 U-234 U-235 U-238 I-125 I-131	* lanized mixing device sampled, treated, and 4.1E-04 5.1E-08 7.1E-09 5.5E-07 7.8E-07 2.8E-08	used to maked a stored. No 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	e homogeneous m releases are assu NA	ixtures of waste med to occur fro NA	e. Solidification a om waste storag	gents are added duri e because the wastes None	ng mixing to transls are fully containe	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13	528						
uilding 5	513 is operated by the filtration Unit filters Stabilization	he Radioactive and Hazardous Wa out waste radioactive particles. Room Air	ste Management Division. The Stabilis In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste	Gross beta zation Unit is a mech waste materials are s H-3 U-234 U-235 U-238 I-125 I-131 Cs-137 C-14	* lanized mixing device sampled, treated, and 4.1E-04 5.1E-08 7.1E-09 5.5E-07 7.8E-07 2.8E-08 2.5E-07 2.6E-04	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	e homogeneous m releases are assu NA	ixtures of waste med to occur fro NA	e. Solidification a om waste storag	gents are added duri e because the wastes None	ng mixing to transls are fully containe	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13 2.5E-12 2.6E-09	528						
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513	Stabilization 1000A	ne Radioactive and Hazardous Wa out waste radioactive particles. Room Air FHE-4	ste Management Division. The Stabili In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste Process optimization and treatibility studies	Gross beta zation Unit is a mech waste materials are s H-3 U-234 U-235 U-238 I-125 I-131 Cs-137 C-14 Cs-134 Ba-133 P-32 Pu-238 Pu-238 Pu-239 Pu-240 Am-241 Th-232 Pu-242 Am-241 Am-243	* lanized mixing device sampled, treated, and 4.1E-04 5.1E-08 7.1E-09 5.5E-07 7.8E-07 2.8E-08 2.5E-07 2.6E-04 3.1E-08 2.0E-09 3.3E-06 1.8E-09 1.4E-07 2.1E-08 4.2E-08 1.3E-06 4.6E-09 7.1E-07	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA 10.5	NA 0.30	NA S.5.5	gents are added duri e because the wastes None HEPA	ng mixing to transles are fully containe 1 0.01	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13 2.5E-12 2.6E-09 3.1E-13 2.0E-14 3.3E-11 1.8E-14 1.4E-12 2.1E-13 4.2E-13 1.3E-11 4.6E-14 7.1E-12	528	NE	8.7E-09	128	SW	1.3E-08	2
513	Stabilization 1000A	ne Radioactive and Hazardous Wa out waste radioactive particles. Room Air FHE-4	ste Management Division. The Stabili. In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste Process optimization and treatibility studies Vacuum filtration of treated	Gross beta zation Unit is a mech waste materials are : H-3 U-234 U-235 U-238 I-125 I-131 Cs-137 C-14 Cs-134 Ba-133 P-32 Pu-238 Pu-239 Pu-240 Am-241 Pu-241 Th-232 Pu-242 Am-241 Am-243 Ba-133	* lanized mixing device sampled, treated, and 4.1E-04 5.1E-08 7.1E-09 5.5E-07 7.8E-07 2.8E-08 2.5E-07 2.6E-04 3.1E-08 2.0E-09 3.3E-06 1.8E-09 1.4E-07 2.1E-08 4.2E-08 1.3E-06 4.6E-09 7.1E-07	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA 10.5	NA 0.30	NA S.5.5	gents are added duri e because the wastes None HEPA	ng mixing to transles are fully containe 1 0.01	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13 2.5E-12 2.6E-09 3.1E-13 3.3E-11 1.8E-14 1.4E-12 2.1E-13 4.2E-13 1.3E-11 4.6E-14 7.1E-12 4.2E-09 1.4E-09 1.1E-09	528	NE	8.7E-09	128	SW	1.3E-08	2
513	Stabilization 1000A	ne Radioactive and Hazardous Wa out waste radioactive particles. Room Air FHE-4	ste Management Division. The Stabili. In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste Process optimization and treatibility studies Vacuum filtration of treated	Gross beta zation Unit is a mech waste materials are : H-3 U-234 U-235 U-238 I-125 I-131 Cs-137 C-14 Cs-134 Ba-133 P-32 Pu-238 Pu-239 Pu-240 Am-241 Pu-241 Th-232 Pu-242 Am-241 Am-243 Ba-133 Be-7	* lanized mixing device sampled, treated, and 4.1E-04 5.1E-08 7.1E-09 5.5E-07 7.8E-07 2.8E-08 2.5E-07 2.6E-04 3.1E-08 2.0E-09 3.3E-06 1.8E-09 1.4E-07 2.1E-08 4.2E-08 4.2E-08 1.3E-06 1.4E-07	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA 10.5	NA 0.30	NA S.5.5	gents are added duri e because the wastes None HEPA	ng mixing to transles are fully containe 1 0.01	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13 2.5E-12 2.6E-09 3.1E-13 2.0E-14 3.3E-11 1.8E-14 1.4E-12 2.1E-13 4.2E-13 1.3E-11 4.6E-14 7.1E-12 4.2E-09 1.4E-09 1.4E-09 1.4E-10	528	NE	8.7E-09	128	SW	1.3E-08	2
silding §	Stabilization 1000A	ne Radioactive and Hazardous Wa out waste radioactive particles. Room Air FHE-4	ste Management Division. The Stabili. In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste Process optimization and treatibility studies Vacuum filtration of treated	Gross beta zation Unit is a mech waste materials are s H-3 U-234 U-235 U-238 I-125 I-131 Cs-137 C-14 Cs-134 Ba-133 P-32 Pu-238 Pu-239 Pu-240 Am-241 Th-232 Pu-241 Th-232 Pu-242 Am-241 Am-243 Ba-133 Be-7 Bi-207	* lanized mixing device sampled, treated, and 4.1E-04 5.1E-08 7.1E-09 5.5E-07 7.8E-07 2.8E-08 2.5E-07 2.6E-04 3.1E-08 2.0E-09 3.3E-06 1.8E-09 1.4E-07 2.1E-08 4.2E-08 1.3E-06 4.6E-09 7.1E-07 4.2E-06 1.4E-07 7.3E-08	1.0E-03	NA 10.5	NA 0.30	NA S.5.5	gents are added duri e because the wastes None HEPA	ng mixing to transles are fully containe 1 0.01	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13 2.5E-12 2.6E-09 3.1E-13 2.0E-14 3.3E-11 1.8E-14 1.4E-12 2.1E-13 4.2E-13 4.2E-13 1.3E-11 4.6E-14 7.1E-12 4.2E-09 1.4E-09 1.4E-09 1.4E-10 7.3E-11	528	NE	8.7E-09	128	SW	1.3E-08	2
silding §	Stabilization 1000A	ne Radioactive and Hazardous Wa out waste radioactive particles. Room Air FHE-4	ste Management Division. The Stabili. In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste Process optimization and treatibility studies Vacuum filtration of treated	Gross beta zation Unit is a mech waste materials are s H-3 U-234 U-235 U-238 I-125 I-131 Cs-137 C-14 Cs-137 C-14 Cs-139 Pu-238 Pu-239 Pu-240 Am-241 Pu-241 Th-232 Pu-242 Am-241 Am-243 Ba-133 Ba-133 Be-7 Bi-207 C-14	* lanized mixing device sampled, treated, and sampled,	1.0E-03	NA 10.5	NA 0.30	NA S.5.5	gents are added duri e because the wastes None HEPA	ng mixing to transles are fully containe 1 0.01	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13 2.5E-12 2.6E-09 3.1E-13 2.0E-14 3.3E-11 1.8E-14 1.4E-12 2.1E-13 4.2E-13 1.3E-11 4.6E-14 7.1E-12 4.2E-09 1.4E-09 1.1E-09 1.4E-10 7.3E-11 1.7E-08	528	NE	8.7E-09	128	SW	1.3E-08	2
silding §	Stabilization 1000A	ne Radioactive and Hazardous Wa out waste radioactive particles. Room Air FHE-4	ste Management Division. The Stabili. In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste Process optimization and treatibility studies Vacuum filtration of treated	Gross beta zation Unit is a mech waste materials are s H-3 U-234 U-235 U-238 I-125 I-131 Cs-137 C-14 Cs-134 Ba-133 P-32 Pu-238 Pu-239 Pu-240 Am-241 Pu-241 Th-232 Pu-242 Am-241 Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109	* lanized mixing device sampled, treated, and 4.1E-04 5.1E-08 7.1E-09 5.5E-07 7.8E-07 2.8E-08 2.5E-07 2.6E-04 3.1E-08 2.0E-09 3.3E-06 1.8E-09 1.4E-07 2.1E-08 4.2E-08 1.3E-06 4.6E-09 7.1E-07 4.2E-06 1.4E-06 1.1E-06 1.4E-07 7.3E-08 1.7E-05 6.9E-10	1.0E-03	NA 10.5	NA 0.30	NA S.5.5	gents are added duri e because the wastes None HEPA	ng mixing to transles are fully containe 1 0.01	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13 2.5E-12 2.6E-09 3.1E-13 2.0E-14 3.3E-11 1.8E-14 1.4E-12 2.1E-13 4.2E-13 1.3E-11 4.6E-14 7.1E-12 4.2E-09 1.4E-09 1.4E-09 1.73E-11 1.7E-08 6.9E-13	528	NE	8.7E-09	128	SW	1.3E-08	2
silding §	Stabilization 1000A	ne Radioactive and Hazardous Wa out waste radioactive particles. Room Air FHE-4	ste Management Division. The Stabili. In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste Process optimization and treatibility studies Vacuum filtration of treated	Gross beta zation Unit is a mech waste materials are s H-3 U-234 U-235 U-238 I-125 I-131 Cs-137 C-14 Cs-137 C-14 Cs-139 Pu-238 Pu-239 Pu-240 Am-241 Pu-241 Th-232 Pu-242 Am-241 Am-243 Ba-133 Ba-133 Be-7 Bi-207 C-14	* lanized mixing device sampled, treated, and sampled,	1.0E-03	NA 10.5	NA 0.30	NA S.5.5	gents are added duri e because the wastes None HEPA	ng mixing to transles are fully containe 1 0.01	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13 2.5E-12 2.6E-09 3.1E-13 2.0E-14 3.3E-11 1.8E-14 1.4E-12 2.1E-13 4.2E-13 1.3E-11 4.6E-14 7.1E-12 4.2E-09 1.4E-09 1.1E-09 1.4E-10 7.3E-11 1.7E-08	528	NE	8.7E-09	128	SW	1.3E-08	2
silding §	Stabilization 1000A	ne Radioactive and Hazardous Wa out waste radioactive particles. Room Air FHE-4	ste Management Division. The Stabili. In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste Process optimization and treatibility studies Vacuum filtration of treated	Gross beta zation Unit is a mech waste materials are s H-3 U-234 U-235 U-238 I-125 I-131 Cs-137 C-14 Cs-134 Ba-133 P-32 Pu-238 Pu-239 Pu-240 Am-241 Pu-241 Th-232 Pu-242 Am-241 Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144	* lanized mixing device sampled, treated, and 4.1E-04 5.1E-08 7.1E-09 5.5E-07 7.8E-07 2.8E-08 2.5E-07 2.6E-04 3.1E-08 2.0E-09 3.3E-06 1.8E-09 1.4E-07 2.1E-08 4.2E-08 1.3E-06 4.6E-09 7.1E-07 4.2E-06 1.4E-06 1.1E-06 1.4E-07 7.3E-08 1.7E-05 6.9E-10 1.1E-05	1.0E-03	NA 10.5	NA 0.30	NA S.5.5	gents are added duri e because the wastes None HEPA	ng mixing to transles are fully containe 1 0.01	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13 2.5E-12 2.6E-09 3.1E-13 3.3E-11 1.8E-14 1.4E-12 2.1E-13 4.2E-13 1.3E-11 4.6E-14 7.1E-12 4.2E-09 1.4E-09 1.1E-09 1.4E-10 7.3E-11 1.7E-08 6.9E-13 1.1E-08	528	NE	8.7E-09	128	SW	1.3E-08	2
silding §	Stabilization 1000A	ne Radioactive and Hazardous Wa out waste radioactive particles. Room Air FHE-4	ste Management Division. The Stabili. In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste Process optimization and treatibility studies Vacuum filtration of treated	Gross beta zation Unit is a mech waste materials are : H-3 U-234 U-235 U-238 I-125 I-131 Cs-137 C-14 Cs-134 Ba-133 P-32 Pu-238 Pu-239 Pu-240 Am-241 Pu-241 Th-232 Pu-242 Am-241 Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244	* lanized mixing device sampled, treated, and 4.1E-04 5.1E-08 7.1E-09 5.5E-07 7.8E-07 2.8E-08 2.5E-07 2.6E-04 3.1E-08 2.0E-09 3.3E-06 1.8E-09 1.4E-07 2.1E-08 4.2E-08 1.3E-06 1.4E-07 7.1E-07	1.0E-03	NA 10.5	NA 0.30	NA S.5.5	gents are added duri e because the wastes None HEPA	ng mixing to transles are fully containe 1 0.01	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13 2.5E-12 2.6E-09 3.1E-13 2.0E-14 3.3E-11 1.8E-14 1.4E-12 2.1E-13 4.2E-13 1.3E-11 4.6E-14 7.1E-12 4.2E-09 1.4E-09 1.4E-09 1.4E-10 7.3E-11 1.7E-08 6.9E-13 1.1E-08 1.3E-09	528	NE	8.7E-09	128	SW	1.3E-08	2
513	Stabilization 1000A	ne Radioactive and Hazardous Wa out waste radioactive particles. Room Air FHE-4	ste Management Division. The Stabili. In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste Process optimization and treatibility studies Vacuum filtration of treated	Gross beta zation Unit is a mech waste materials are s H-3 U-234 U-235 U-238 I-125 I-131 Cs-137 C-14 Cs-134 Ba-133 P-32 Pu-238 Pu-239 Pu-240 Am-241 Pu-241 Th-232 Pu-242 Am-241 Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Cm-244 Co-57 Co-58 Co-60	* lanized mixing device sampled, treated, and sampled,	1.0E-03	NA 10.5	NA 0.30	NA S.5.5	gents are added duri e because the wastes None HEPA	ng mixing to transles are fully containe 1 0.01	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13 2.5E-12 2.6E-09 3.1E-13 2.0E-14 3.3E-11 1.8E-14 1.4E-12 2.1E-13 4.2E-13 1.3E-11 4.6E-14 7.1E-12 4.2E-09 1.4E-09 1.1E-09 1.4E-10 7.3E-11 1.7E-08 6.9E-13 1.1E-08 1.3E-09 2.2E-10 1.4E-10 9.1E-10	528	NE	8.7E-09	128	SW	1.3E-08	2
513	Stabilization 1000A	ne Radioactive and Hazardous Wa out waste radioactive particles. Room Air FHE-4	ste Management Division. The Stabili. In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste Process optimization and treatibility studies Vacuum filtration of treated	Gross beta zation Unit is a mech waste materials are s H-3 U-234 U-235 U-238 I-125 I-131 CS-137 C-14 CS-134 Ba-133 P-32 PU-238 PU-239 PU-240 Am-241 Th-232 PU-242 Am-241 Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51	* lanized mixing device sampled, treated, and sampled,	1.0E-03	NA 10.5	NA 0.30	NA S.5.5	gents are added duri e because the wastes None HEPA	ng mixing to transles are fully containe 1 0.01	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13 2.5E-12 2.6E-09 3.1E-13 2.0E-14 3.3E-11 1.8E-14 1.4E-12 2.1E-13 4.2E-13 1.3E-11 4.6E-14 7.1E-12 4.2E-09 1.1E-09 1.4E-09 1.1E-09 1.4E-10 7.3E-11 1.7E-08 6.9E-13 1.1E-08 1.3E-09 2.2E-10 1.4E-10 9.1E-10	528	NE	8.7E-09	128	SW	1.3E-08	2
ne Micro	Stabilization 1000A	ne Radioactive and Hazardous Wa out waste radioactive particles. Room Air FHE-4	ste Management Division. The Stabili. In the Laboratory, small quantities of Treatment of hazardous, mixed or radioactive waste Process optimization and treatibility studies Vacuum filtration of treated	Gross beta zation Unit is a mech waste materials are s H-3 U-234 U-235 U-238 I-125 I-131 Cs-137 C-14 Cs-134 Ba-133 P-32 Pu-238 Pu-239 Pu-240 Am-241 Pu-241 Th-232 Pu-242 Am-241 Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Cm-244 Co-57 Co-58 Co-60	* lanized mixing device sampled, treated, and sampled,	1.0E-03	NA 10.5	NA 0.30	NA S.5.5	gents are added duri e because the wastes None HEPA	ng mixing to transles are fully containe 1 0.01	4.1E-07 5.1E-11 7.1E-12 5.5E-10 7.8E-12 2.8E-13 2.5E-12 2.6E-09 3.1E-13 2.0E-14 3.3E-11 1.8E-14 1.4E-12 2.1E-13 4.2E-13 1.3E-11 4.6E-14 7.1E-12 4.2E-09 1.4E-09 1.1E-09 1.4E-10 7.3E-11 1.7E-08 6.9E-13 1.1E-08 1.3E-09 2.2E-10 1.4E-10 9.1E-10	528	NE	8.7E-09	128	SW	1.3E-08	2

	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated	10 mrem/y S	<u> Site-Wide Dos</u>	se Requirement	<u>0.1 mren</u>	n/y Monitoring	Requirement	Source
					with Potential for	State	Height (m)	Diameter	Velocity	Device(s)		Annual Emissions			EDE	Distance	Direction	Unabated	Category
					Release (Ci)	Factor		(m)	(m/s)		Factor	(Ci)	SWMEI (m)	to SWMEI	(mrem)	to MEI (m)	to MEI	EDE (mrem)	
514	108	(continued)		Eu-152	4.2E-06	1.0E-03						4.2E-09							
				Eu-154	4.2E-06	1.0E-03						4.2E-09							
				Eu-155	3.3E-07	1.0E-03						3.3E-10							
				Fe-55	1.4E-07	1.0E-03						1.4E-10							
				Gd-148	1.1E-06	1.0E-03						1.1E-09							
				H-3	1.2E-04	1.0E-03						1.2E-07							+
				Hf-172	1.4E-07	1.0E-03						1.4E-10				+			+
				K-40	7.0E-10	1.0E-03						7.0E-13							
				Lu-174	1.4E-07	1.0E-03						1.4E-10 1.4E-13							+
				MFP	1.4E-10	1.0E-03													+
				Mn-54	3.5E-07	1.0E-03						3.5E-10							
				Na-22 Np-237	2.1E-05 1.1E-06	1.0E-03 1.0E-03						2.1E-08 1.1E-09						·	+
				P-32	1.1E-06 1.2E-03	1.0E-03						1.2E-06							
				Pb-210	1.2E-05	1.0E-03						1.2E-09							+
				Pu-238	2.8E-06	1.0E-03						2.8E-09	-			+			+
				Pu-239	1.1E-05	1.0E-03						1.1E-08				+			+
				Pu-240	3.3E-08	1.0E-03						3.3E-11				+			+
				Pu-241	1.5E-07	1.0E-03						1.5E-10				+			_
				Pu-241 Pu-242	1.1E-06	1.0E-03					+	1.1E-09						<u> </u>	+
				Pu-242 Pu-244	2.8E-08	1.0E-03		+			+	2.8E-11	 	+ +					+
				Ra-226	9.1E-08	1.0E-03		1	+			9.1E-11	+						+
				Sb-125	3.3E-07	1.0E-03						3.3E-10						<u>'</u>	+
				Sc-46	1.4E-07	1.0E-03						1.4E-10						<u> </u>	+
				Sr-90	1.7E-09	1.0E-03						1.7E-12							+
				Tc-99	1.1E-06	1.0E-03						1.1E-09							
				Th-228	2.0E-10	1.0E-03						2.0E-13						·	1
				Th-229	1.1E-06	1.0E-03						1.1E-09						<u>'</u>	+
				Th-230	1.1E-06	1.0E-03						1.1E-09							+
				Th-232	1.3E-06	1.0E-03						1.3E-09							
				U-232	1.1E-06	1.0E-03						1.1E-09							+
				U-233	1.1E-06	1.0E-03						1.1E-09						<u></u>	
				U-234	2.1E-05	1.0E-03						2.1E-08							+
				U-235	1.5E-06	1.0E-03						1.5E-09							_
				U-238	8.2E-05	1.0E-03						8.2E-08							+
				Y-88	2.7E-10	1.0E-03						2.7E-13							
514	Evaporator	Room Air	Waste consolidation	Am-241	4.5E-05	1.0E-03	NA	NA	NA	None	1	4.5E-08	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-241 Am-243	4.5E-05 1.5E-05	1.0E-03 1.0E-03	NA	NA	NA	None	1	4.5E-08 1.5E-08	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation		1.5E-05 1.2E-05		NA	NA	NA	None	1	1.5E-08 1.2E-08	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243	1.5E-05	1.0E-03	NA	NA	NA	None	1	1.5E-08	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133	1.5E-05 1.2E-05	1.0E-03 1.0E-03	NA	NA	NA	None	1	1.5E-08 1.2E-08	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7	1.5E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04	1.0E-03 1.0E-03 1.0E-03	NA	NA	NA	None	1	1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207	1.5E-05 1.2E-05 1.5E-06 7.8E-07	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA	NA	NA NA	None	1	1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14	1.5E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA	NA	NA	None	1	1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109	1.5E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA	NA	None	1	1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57	1.5E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA	NA NA	None	1	1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58	1.5E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA	NA NA	None	1	1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60	1.5E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA NA	NA NA	None	1	1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 9.7E-09	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51	1.5E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA NA	NA	None	1	1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 9.7E-09 1.5E-09	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134	1.5E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06 1.5E-06 1.2E-05	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA	NA	None	1	1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 9.7E-09 1.2E-08	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137	1.5E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06 1.5E-06 1.2E-05 7.5E-05	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA NA	NA	None	1	1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 9.7E-09 9.7E-09 1.5E-09 1.2E-08 7.5E-08	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152	1.5E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06 1.5E-06 1.2E-05 7.5E-05 4.5E-05	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA NA	NA	None	1	1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 9.7E-09 1.2E-08 7.5E-08	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154	1.5E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06 1.5E-06 1.2E-05 7.5E-05 4.5E-05	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA NA	NA	None	1	1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 9.7E-09 1.5E-09 1.5E-08 4.5E-08	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155	1.5E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06 1.5E-05 1.5E-05 4.5E-05 4.5E-05 3.5E-05	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA NA	NA NA	None	1	1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-08 4.5E-08 4.5E-08 3.5E-09	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55	1.5E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06 1.5E-05 4.5E-05 4.5E-05 4.5E-05 3.5E-06 1.5E-06	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA NA	NA	None	1	1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 9.7E-09 1.2E-08 7.5E-08 4.5E-08 4.5E-08 3.5E-09 1.5E-09	528	NE	1.2E-03	217	SW	3.1E-03	1
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55 Gd-148	1.5E-05 1.2E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 1.5E-06 1.2E-05 7.5E-05 4.5E-05 3.5E-06 1.5E-06 1.5E-06	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA NA	NA	None	1	1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 9.7E-09 1.5E-09 1.2E-08 4.5E-08 4.5E-08 4.5E-08 1.5E-09 1.5E-09	528	NE	1.2E-03	217	SW	3.1E-03	
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55 Gd-148 H-3	1.5E-05 1.2E-05 1.2E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06 1.5E-05 7.5E-05 4.5E-05 4.5E-05 1.5E-06 1.5E-06 1.5E-06	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA NA	NA NA	None		1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 9.7E-09 1.5E-09 1.5E-09 4.5E-08 4.5E-08 4.5E-08 3.5E-09 1.2E-08 1.2E-08	528	NE	1.2E-03	217	SW	3.1E-03	
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55 Gd-148 H-3 Hf-172	1.5E-05 1.2E-05 1.2E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06 1.5E-05 4.5E-05 4.5E-05 3.5E-06 1.2E-05 1.5E-06 1.5E-06	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA NA	NA NA	None		1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 9.7E-09 1.2E-08 7.5E-08 4.5E-08 4.5E-08 3.5E-09 1.2E-08 1.2E-08	528	NE	1.2E-03	217	SW	3.1E-03	
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55 Gd-148 H-3 Hf-172 K-40	1.5E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06 1.5E-05 4.5E-05 4.5E-05 4.5E-05 1.2E-05 1.2E-05 1.2E-06 1.2E-06	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA NA	NA NA	None		1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 9.7E-09 1.5E-09 1.5E-09 1.5E-08 4.5E-08 4.5E-08 3.5E-09 1.2E-08 1.5E-09 1.5E-09	528	NE	1.2E-03	217	SW	3.1E-03	
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-155 Fe-55 Gd-148 H-3 Hf-172 K-40 Lu-174	1.5E-05 1.2E-05 1.2E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06 1.5E-05 4.5E-05 4.5E-05 3.5E-06 1.2E-05 1.2E-05 7.5E-06 1.5E-06 1.5E-06 1.5E-06	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA NA	NA NA	None		1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 1.5E-09 1.2E-08 7.5E-08 4.5E-08 4.5E-08 1.5E-09 1.2E-08 1.5E-09 1.5E-09	528	NE	1.2E-03	217	SW	3.1E-03	
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55 Gd-148 H-3 Hf-172 K-40 Lu-174 MFP	1.5E-05 1.2E-05 1.2E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 1.5E-06 1.2E-05 7.5E-05 4.5E-05 3.5E-06 1.2E-05 1.2E-03 1.5E-06 1.2E-03 1.5E-06 1.5E-06 1.2E-03 1.5E-06 1.5E-06 1.5E-06	1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03 1.0E-03	NA NA	NA NA	NA NA	None		1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 1.5E-09 1.2E-08 4.5E-08 4.5E-08 4.5E-08 1.5E-09 1.2E-06 1.5E-09 1.2E-06 1.5E-09	528	NE	1.2E-03	217	SW	3.1E-03	
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55 Gd-148 H-3 Hf-172 K-40 Lu-174 MFP Mn-54	1.5E-05 1.2E-05 1.2E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 1.5E-06 1.5E-05 4.5E-05 4.5E-05 1.2E-03 1.5E-06 1.2E-03 1.5E-06 1.5E-06 1.5E-06 1.5E-09 1.5E-09 1.5E-09	1.0E-03 1.0E-03	NA NA	NA NA	NA NA	None		1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 1.5E-09 1.2E-08 4.5E-08 4.5E-08 4.5E-08 1.2E-06 1.5E-09 1.2E-08 1.2E-06 1.5E-09 1.2E-08	528	NE	1.2E-03	217	SW	3.1E-03	
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55 Gd-148 H-3 Hf-172 K-40 Lu-174 MFP Mn-54 Na-22	1.5E-05 1.2E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 1.5E-06 1.5E-05 4.5E-05 4.5E-05 1.2E-05 1.2E-05 1.5E-06	1.0E-03 1.0E-03	NA NA	NA NA	NA NA	None		1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-08 4.5E-08 4.5E-08 3.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09	528	NE	1.2E-03	217	SW	3.1E-03	
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55 Gd-148 H-3 Hf-172 K-40 Lu-174 MFP Mn-54 Na-22 Np-237	1.5E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06 1.5E-05 7.5E-05 4.5E-05 4.5E-05 1.2E-05 1.2E-05 1.2E-06 1.5E-06 1.5E-09 1.5E-06	1.0E-03 1.0E-03	NA NA	NA NA	NA NA	None		1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 1.5E-09 1.2E-08 7.5E-08 4.5E-08 3.5E-09 1.2E-08 1.2E-08 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09	528	NE	1.2E-03	217	SW	3.1E-03	
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55 Gd-148 H-3 Hf-172 K-40 Lu-174 MFP Mn-54 Na-22 Np-237 P-32	1.5E-05 1.2E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06 1.5E-05 4.5E-05 4.5E-05 1.2E-05 1.2E-03 1.5E-06 1.2E-03 1.5E-06 1.2E-03 1.5E-06 1.2E-03 1.5E-06 1.2E-05 1.2E-05 1.2E-03	1.0E-03 1.0E-03	NA NA	NA NA	NA NA	None		1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 1.5E-09 1.5E-09 1.2E-08 4.5E-08 4.5E-08 3.5E-09 1.2E-08 1.2E-08 1.2E-08 1.2E-08 1.2E-08 1.2E-08 1.2E-08 1.2E-08 1.2E-08	528	NE	1.2E-03	217	SW	3.1E-03	
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55 Gd-148 H-3 Hf-172 K-40 Lu-174 MFP Mn-54 Na-22 Np-237 P-32 Pb-210	1.5E-05 1.2E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06 1.5E-05 4.5E-05 4.5E-05 1.2E-05 1.2E-05 1.2E-05 1.2E-06 1.2E-05	1.0E-03 1.0E-03	NA NA	NA NA	NA NA	None		1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 1.5E-09 1.2E-08 4.5E-08 4.5E-08 4.5E-08 1.2E-06 1.5E-09 1.2E-06 1.5E-09 1.2E-08 1.2E-06 1.5E-09 1.2E-08 1.2E-06 1.3E-09	528	NE	1.2E-03	217	SW	3.1E-03	
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55 Gd-148 H-3 Hf-172 K-40 Lu-174 MFP Mn-54 Na-22 Np-237 P-32 Pb-210 Pu-238	1.5E-05 1.2E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 1.5E-06 1.5E-05 4.5E-05 4.5E-05 1.2E-03 1.5E-06 1.2E-03 1.5E-06 1.5E-06 1.2E-03 1.5E-06 1.5E-06 1.2E-03 1.5E-06 1.5E-06 1.5E-06 1.2E-05 1.2E-03 1.5E-06 1.5E-06 1.5E-06 1.5E-06 1.5E-06 1.5E-09 3.7E-06 1.5E-09 3.7E-06 1.5E-09 3.7E-06 1.5E-09 3.7E-06 1.5E-05 3.7E-06 1.5E-05	1.0E-03 1.0E-03	NA NA	NA NA	NA NA	None		1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 1.5E-09 1.2E-08 4.5E-08 4.5E-08 4.5E-08 1.2E-06 1.5E-09 1.2E-06 1.5E-09 1.2E-08 1.2E-06 1.5E-09 1.2E-08 1.2E-06 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-09	528	NE	1.2E-03	217	SW	3.1E-03	
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55 Gd-148 H-3 Hf-172 K-40 Lu-174 MFP Mn-54 Na-22 Np-237 P-32 Pb-210 Pu-238 Pu-239	1.5E-05 1.2E-05 1.2E-05 1.5E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 1.5E-06 1.2E-05 4.5E-05 4.5E-05 1.2E-03 1.5E-06 1.5E-06 1.5E-06 1.2E-03 1.5E-06 1.2E-03 1.5E-06 1.5E-06 1.5E-06 1.2E-05 1.2E-03 1.5E-06 1.5E-06 1.5E-06 1.5E-09 3.7E-06 1.5E-09 3.7E-06 2.6E-06 1.2E-05 6.4E-03 1.3E-05 3.0E-05	1.0E-03	NA NA	NA NA	NA NA	None		1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 1.5E-09 1.2E-08 4.5E-08 4.5E-08 4.5E-08 1.2E-06 1.5E-09 1.2E-08 1.2E-06 1.5E-09 1.2E-08 1.2E-06 1.5E-09 1.3E-09	528	NE	1.2E-03	217	SW	3.1E-03	
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55 Gd-148 H-3 Hf-172 K-40 Lu-174 MFP Mn-54 Na-22 Np-237 P-32 Pb-210 Pu-238 Pu-239 Pu-240	1.5E-05 1.2E-05 1.2E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 1.5E-06 1.5E-06 1.2E-05 4.5E-05 4.5E-05 1.2E-03 1.5E-06 1.5E-06 1.5E-06 1.2E-05 1.2E-03 1.5E-06 1.5E-06 1.2E-05 1.2E-03 1.5E-06 1.5E-06 1.5E-06 1.5E-06 1.5E-06 1.5E-09 1.5E-06 1.5E-09 1.5E-06 1.5E-09 3.7E-06 2.6E-06 1.2E-05 6.4E-03 1.3E-05 6.4E-03 1.3E-05 3.0E-05	1.0E-03	NA NA	NA NA	NA NA	None		1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 9.7E-09 1.5E-09 1.2E-08 4.5E-08 4.5E-08 4.5E-08 1.2E-06 1.5E-09 1.5E-09 1.2E-08 1.2E-06 1.5E-09 1.5E-12 1.5E-09	528	NE	1.2E-03	217	SW	3.1E-03	
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55 Gd-148 H-3 Hf-172 K-40 Lu-174 MFP Mn-54 Na-22 Np-237 P-32 Pb-210 Pu-238 Pu-239 Pu-240 Pu-241	1.5E-05 1.2E-05 1.2E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 9.7E-06 1.5E-05 4.5E-05 4.5E-05 1.2E-05 1.2E-05 1.2E-05 1.2E-06 1.5E-06 1.2E-05 1.2E-06 1.5E-09 1.5E-06 1.5E-09 1.5E-06 1.5E-09 1.5E-06 1.5E-09 1.5E-06 1.2E-05 1.2E-05 1.2E-05 1.2E-05 1.2E-05	1.0E-03	NA NA	NA NA	NA NA	None		1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 1.5E-09 1.5E-09 1.5E-08 4.5E-08 3.5E-09 1.5E-09 1.5E-12 3.7E-09 1.5E-12 3.7E-09 1.6E-09 1.5E-12 3.7E-09	528	NE	1.2E-03	217	SW	3.1E-03	
514	Evaporator	Room Air	Waste consolidation	Am-243 Ba-133 Be-7 Bi-207 C-14 Cd-109 Ce-144 Cm-244 Co-57 Co-58 Co-60 Cr-51 Cs-134 Cs-137 Eu-152 Eu-154 Eu-155 Fe-55 Gd-148 H-3 Hf-172 K-40 Lu-174 MFP Mn-54 Na-22 Np-237 P-32 Pb-210 Pu-238 Pu-239 Pu-240	1.5E-05 1.2E-05 1.2E-06 7.8E-07 1.8E-04 7.4E-09 1.2E-04 1.4E-05 2.3E-06 1.5E-06 1.5E-06 1.5E-06 1.2E-05 4.5E-05 4.5E-05 1.2E-03 1.5E-06 1.5E-06 1.5E-06 1.2E-05 1.2E-03 1.5E-06 1.5E-06 1.2E-05 1.2E-03 1.5E-06 1.5E-06 1.5E-06 1.5E-06 1.5E-06 1.5E-09 1.5E-06 1.5E-09 1.5E-06 1.5E-09 3.7E-06 2.6E-06 1.2E-05 6.4E-03 1.3E-05 6.4E-03 1.3E-05 3.0E-05	1.0E-03	NA NA	NA NA	NA NA	None		1.5E-08 1.2E-08 1.5E-09 7.8E-10 1.8E-07 7.4E-12 1.2E-07 1.4E-08 2.3E-09 1.5E-09 9.7E-09 1.5E-09 1.2E-08 4.5E-08 4.5E-08 4.5E-08 1.2E-06 1.5E-09 1.5E-09 1.2E-08 1.2E-06 1.5E-09 1.5E-12 1.5E-09	528	NE	1.2E-03	217	SW	3.1E-03	

Building Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated			ose Requirement		/y Monitoring		Source
				with Potential for	State	Height (m)	Diameter	Velocity	Device(s)		Annual Emissions	Distance to			Distance	Direction	Unabated	Categor
514 Evaporator	(continued)		Sb-125	Release (Ci) 3.5E-06	Factor 1.0E-03		(m)	(m/s)		Factor	(Ci) 3.5E-09	SWMEI (m)	to SWMEI	(mrem)	to MEI (m)	to MEI	EDE (mrem)	+
514 Evaporator	(continued)		Sc-46	1.5E-06	1.0E-03						1.5E-09							+
			Sr-90	1.8E-08	1.0E-03						1.8E-11							+
			Tc-99	1.2E-05	1.0E-03						1.2E-08							+
			Th-228	2.1E-09	1.0E-03						2.1E-12							
			Th-229	1.2E-05	1.0E-03						1.2E-08							
			Th-230	1.2E-05	1.0E-03						1.2E-08							
			Th-232	1.3E-05	1.0E-03						1.3E-08							
			U-232	1.2E-05	1.0E-03						1.2E-08							
			U-233	1.2E-05	1.0E-03						1.2E-08							
			U-234	2.3E-04	1.0E-03						2.3E-07							_
			U-235	1.6E-05	1.0E-03						1.6E-08							
			U-238 Y-88	8.8E-04 2.9E-09	1.0E-03 1.0E-03						8.8E-07 2.9E-12							_
			1-00	2.9E-09	1.02-03						2.9E-12							+
Building 612 is operated by the R	adioactive and Hazardous Wa	aste Management Division. It is a fa	 cility in which waste is	renackaged for shipm	lent off site													+
612 100	Room Air	Waste sampling	Am-241	6.3E-08	1.0E-06	NA	NA	NA	None	1	6.3E-14	444	NNE	1.1E-06	276	SW	2.3E-06	2
0.12	11001117111	Trace camping	Am-243	5.3E-13	1.0E-06		101		. 10.10		5.3E-19		11112	2 00			2.02.00	+
			Au-195	3.1E-12	1.0E-06						3.1E-18							1
			Ba-133	1.3E-10	1.0E-06						1.3E-16							
			Ba-140	4.2E-09	1.0E-06						4.2E-15							
			Be-7	1.8E-09	1.0E-06						1.8E-15							
			C-14	1.0E-04	1.0E-06						1.0E-10			1		1		
			Cd-109	2.5E-12	1.0E-06						2.5E-18					1		
			Ce-141	1.6E-08	1.0E-06			+		+	1.6E-14 1.4E-14			-		1		+
			Ce-144 Cf-250	1.4E-08 9.1E-13	1.0E-06 1.0E-06						9.1E-19					-		_
			CI-230	2.1E-11	1.0E-06						2.1E-17							_
			Co-60	2.1E-08	1.0E-06						2.1E-14							+
			Cr-51	9.1E-10	1.0E-06						9.1E-16							+
			Cs-134	2.0E-09	1.0E-06						2.0E-15							+
			Cs-137	4.5E-08	1.0E-06						4.5E-14							
			Eu-152	9.1E-10	1.0E-06						9.1E-16							
			Eu-154	9.1E-10	1.0E-06						9.1E-16							
			Eu-155	9.2E-10	1.0E-06						9.2E-16							
			Fe-55	5.5E-09	1.0E-06						5.5E-15							
			H-3	2.3E-03	1.0E-06						2.3E-09							
			I-125	5.2E-08	1.0E-06						5.2E-14							+
			I-131 K-40	1.8E-09 7.7E-10	1.0E-06 1.0E-06						1.8E-15 7.7E-16							+
			Mn-54	1.2E-10	1.0E-06			+			1.2E-16							
			Mo-99	8.9E-10	1.0E-06						8.9E-16							+
			Nb-94	9.1E-10	1.0E-06						9.1E-16							+
			Nb-95	1.4E-08	1.0E-06						1.4E-14							1
			Nd-147	9.1E-10	1.0E-06						9.1E-16							
			Np-237	4.0E-15	1.0E-06						4.0E-21							
			Np-239	7.3E-10	1.0E-06						7.3E-16							
			P-32	1.7E-05	1.0E-06						1.7E-11							
			Pm-147	8.2E-11	1.0E-06						8.2E-17							
			Pm-151	2.7E-10	1.0E-06			1			2.7E-16			-		1		
			Pt-195m Pu-238	5.5E-10 4.2E-09	1.0E-06 1.0E-06						5.5E-16 4.2E-15			-	+			_
			Pu-238 Pu-239	2.5E-08	1.0E-06			+			2.5E-14			 		1		+
			Pu-240	1.4E-09	1.0E-06			+			1.4E-15			<u> </u>				+
			Pu-240	3.3E-08	1.0E-06						3.3E-14							+
			Pu-242	2.8E-08	1.0E-06						2.8E-14							1
			Ra-223	2.7E-12	1.0E-06						2.7E-18							1
			Ra-226	1.3E-12	1.0E-06						1.3E-18							
			Rh-103	9.1E-09	1.0E-06						9.1E-15							
			Ru-106	1.4E-08	1.0E-06						1.4E-14							
			S-35	3.9E-06	1.0E-06						3.9E-12			-	1	1		
			Sb-125	4.4E-12	1.0E-06						4.4E-18				1	1		_
			Sm-151	8.2E-12	1.0E-06						8.2E-18			-	-			+
			Sr-90	7.1E-09	1.0E-06			+			7.1E-15			+		-		+
			Th-228 Th-230	7.8E-10 6.8E-10	1.0E-06 1.0E-06			+		+	7.8E-16 6.8E-16			+		+		+
			Th-232	1.2E-09	1.0E-06 1.0E-06						1.2E-15			1		1		+
			U-233	1.2E-09 1.8E-09	1.0E-06			+			1.8E-15			 				+
			U-234	6.4E-05	1.0E-06			+		+	6.4E-11			+		+		+
			U-235	2.8E-06	1.0E-06						2.8E-12					1		+
			U-237	1.9E-08	1.0E-06						1.9E-14							+
			U-238	6.0E-05	1.0E-06						6.0E-11							+
			Zr-95	3.4E-08	1.0E-06						3.4E-14							\top
			MFP	2.1E-08	1.0E-06						2.1E-14							\top

Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated	10 mrem/y S	Site-Wide Do	se Requirement	0.1 mrem	/y Monitoring	Requirement	Source
					with Potential for	State	Height (m)	Diameter	Velocity	Device(s)	Abatement	Annual Emissions			EDE	Distance	Direction	Unabated	Category
					Release (Ci)	Factor		(m)	(m/s)		Factor	(Ci)	SWMEI (m)	to SWMEI	(mrem)	to MEI (m)	to MEI	EDE (mrem)	
612	101	FHE-4	Laboratory analysis	Am-241	4.2E-05	1.0E-03	10.5	0.31	5.6	HEPA	0.01	4.2E-10 1.4E-10	444	NE	8.3E-06	295	ENE	1.0E-03	1
			of waste treatment	Am-243 Ba-133	1.4E-05 1.1E-05	1.0E-03 1.0E-03						1.4E-10 1.1E-10							+
			and treatability samples	Ba-133 Be-7	1.1E-05 1.4E-06	1.0E-03 1.0E-03						1.4E-11							+
				Bi-207	7.3E-07	1.0E-03						7.3E-12							
				C-14	1.7E-04	1.0E-03						1.7E-09							
				Cd-109	6.9E-09	1.0E-03						6.9E-14							
				Ce-144	1.1E-04	1.0E-03						1.1E-09							
				Cm-244	1.3E-05	1.0E-03						1.3E-10							
				Co-57	2.2E-06	1.0E-03						2.2E-11							
				Co-58	1.4E-06	1.0E-03						1.4E-11							
				Co-60	9.1E-06	1.0E-03						9.1E-11							
				Cr-51	1.4E-06	1.0E-03						1.4E-11							
				Cs-134	1.1E-05	1.0E-03						1.1E-10							
				Cs-137	7.0E-05	1.0E-03						7.0E-10							
				Eu-152 Eu-154	4.2E-05 4.2E-05	1.0E-03 1.0E-03						4.2E-10 4.2E-10							
				Eu-155	3.3E-06	1.0E-03						3.3E-11							+
	+			Fe-55	1.4E-06	1.0E-03					+	1.4E-11					+ -		
				Gd-148	1.1E-05	1.0E-03						1.1E-10							
				H-3	1.2E-03	1.0E-03						1.2E-08							
				Hf-172	1.4E-06	1.0E-03						1.4E-11							
				K-40	7.0E-09	1.0E-03						7.0E-14							
				Lu-174	1.4E-06	1.0E-03						1.4E-11							
				MFP	1.4E-09	1.0E-03						1.4E-14							
				Mn-54	3.5E-06	1.0E-03						3.5E-11							1
				Na-22	2.1E-04	1.0E-03						2.1E-09							
				Np-237	1.1E-05	1.0E-03						1.1E-10							
				P-32 Pb-210	1.2E-02 1.2E-05	1.0E-03 1.0E-03						1.2E-07 1.2E-10							
				Pu-238	2.8E-05	1.0E-03						2.8E-10							_
				Pu-239	1.1E-04	1.0E-03						1.1E-09							
				Pu-240	3.3E-07	1.0E-03						3.3E-12							
				Pu-241	1.5E-06	1.0E-03						1.5E-11							
				Pu-242	1.1E-05	1.0E-03						1.1E-10							
				Pu-244	2.8E-07	1.0E-03						2.8E-12							
				Ra-226	9.1E-07	1.0E-03						9.1E-12							
				Sb-125	3.3E-06	1.0E-03						3.3E-11							
				Sc-46	1.4E-06	1.0E-03						1.4E-11							
				Sr-90	1.7E-08	1.0E-03						1.7E-13							
				Tc-99	1.1E-05	1.0E-03						1.1E-10							
				Th-228	2.0E-09	1.0E-03						2.0E-14 1.1E-10							+
				Th-229 Th-230	1.1E-05 1.1E-05	1.0E-03 1.0E-03						1.1E-10 1.1E-10							
				Th-232	1.3E-05	1.0E-03						1.3E-10							
				U-232	1.1E-05	1.0E-03						1.1E-10							
				U-233	1.1E-05	1.0E-03						1.1E-10							
				U-234	2.1E-04	1.0E-03						2.1E-09							
				U-235	1.5E-05	1.0E-03						1.5E-10							
				U-238	8.2E-04	1.0E-03						8.2E-09							
		<u> </u>		Y-88	2.7E-09	1.0E-03				_		2.7E-14							
612	102	Room Air	Laboratory analysis	Am-241	4.2E-05	1.0E-03	NA	NA	NA	None	1	4.2E-08	444	NE	1.1E-03	295	ENE	1.5E-03	1 1
			of waste treatment	Am-243	1.4E-05	1.0E-03						1.4E-08	-						
			and treatability samples	Ba-133	1.1E-05	1.0E-03		-				1.1E-08							
				Be-7 Bi-207	1.4E-06 7.3E-07	1.0E-03 1.0E-03		-				1.4E-09 7.3E-10							
				C-14	1.7E-04	1.0E-03						1.7E-07							
				Cd-109	6.9E-09	1.0E-03						6.9E-12							
				Ce-144	1.1E-04	1.0E-03						1.1E-07							
				Cm-244	1.3E-05	1.0E-03						1.3E-08							
				Co-57	2.2E-06	1.0E-03						2.2E-09							
				Co-58	1.4E-06	1.0E-03						1.4E-09							
				Co-60	9.1E-06	1.0E-03						9.1E-09							
	·			Cr-51	1.4E-06	1.0E-03						1.4E-09							
				Cs-134	1.1E-05	1.0E-03						1.1E-08							
				Cs-137	7.0E-05	1.0E-03		-				7.0E-08							
				Eu-152	4.2E-05	1.0E-03						4.2E-08							1
				Eu-154	4.2E-05	1.0E-03						4.2E-08							
\vdash				Eu-155	3.3E-06	1.0E-03						3.3E-09							
				Fe-55 Gd-148	1.4E-06 1.1E-05	1.0E-03 1.0E-03						1.4E-09 1.1E-08							-
 				H-3	1.1E-05 1.2E-03	1.0E-03 1.0E-03						1.1E-08 1.2E-06							-
				_ п-3	1.4E-U3	1.05-03		I				1.46-06							

Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated	10 mrem/y	Site-Wide Do	se Requirement	() 1 mrem/	y Monitoring	Requirement	Source
1	ROOM) Area	Stack ID	Орстаноп	Radionaciaes	with Potential for	State	Height (m)	Diameter	Velocity	Device(s)	Abatement	Annual Emissions			EDE EDE	Distance	Direction		Category
					Release (Ci)	Factor	Transpire (m)	(m)	(m/s)		Factor	(Ci)	SWMEI (m)		(mrem)	to MEI (m)	to MEI	EDE (mrem)	- carrigery
612	102	(continued)		Hf-172	1.4E-06	1.0E-03		, ,	, ,			1.4E-09	· · · ·		,	, ,		` '	
				K-40	7.0E-09	1.0E-03						7.0E-12							
				Lu-174	1.4E-06	1.0E-03						1.4E-09							
				MFP	1.4E-09	1.0E-03						1.4E-12							
				Mn-54	3.5E-06	1.0E-03						3.5E-09							
				Na-22	2.1E-04	1.0E-03						2.1E-07							
				Np-237	1.1E-05	1.0E-03						1.1E-08							
				P-32	1.2E-02	1.0E-03						1.2E-05							
				Pb-210	1.2E-05	1.0E-03						1.2E-08							
				Pu-238	2.8E-05	1.0E-03						2.8E-08							
				Pu-239	1.1E-04	1.0E-03						1.1E-07							
				Pu-240	3.3E-07	1.0E-03						3.3E-10							
				Pu-241	1.5E-06	1.0E-03						1.5E-09							
				Pu-242	1.1E-05	1.0E-03						1.1E-08							
				Pu-244	2.8E-07	1.0E-03						2.8E-10							
				Ra-226	9.1E-07	1.0E-03						9.1E-10							
				Sb-125	3.3E-06	1.0E-03						3.3E-09							
				Sc-46	1.4E-06	1.0E-03						1.4E-09							
				Sr-90	1.7E-08	1.0E-03						1.7E-11							
				Tc-99	1.1E-05	1.0E-03						1.1E-08	1						
				Th-228	2.0E-09	1.0E-03						2.0E-12	1						
				Th-229	1.1E-05	1.0E-03						1.1E-08							
<u> </u>				Th-230	1.1E-05	1.0E-03	-		1		1	1.1E-08	1						
<u> </u>				Th-232	1.3E-05	1.0E-03			1		1	1.3E-08	1						
				U-232	1.1E-05	1.0E-03						1.1E-08							
				U-233	1.1E-05	1.0E-03	-					1.1E-08							
				U-234	2.1E-04	1.0E-03						2.1E-07	1						
				U-235	1.5E-05	1.0E-03						1.5E-08							
				U-238	8.2E-04	1.0E-03						8.2E-07							
				Y-88	2.7E-09	1.0E-03						2.7E-12							
Building 625	5 is operated by Ra	adioactive and Hazardous Waste	Management.																
625	Repack Tent	FHE	Waste inspection and repackaging	Am-241	8.8E-09	1.0E-06	1.5	0.31	6.9	HEPA	0.01	8.8E-17	378	NE	6.6E-13	289	ENE	8.2E-11	1
-				Ba-133	1.1E-10	1.0E-06						1.1E-18							
				Ce-144	1.0E-13	1.0E-06						1.0E-21							
				Cm-243	1.2E-11	1.0E-06						1.2E-19							
				Cm-245	1.7E-12	1.0E-06						1.7E-20							
				Co-60	1.4E-10	1.0E-06						1.4E-18							
				Cs-137	2.8E-09	1.0E-06						2.8E-17							
				H-3	1.0E-08	1.0E-06						1.0E-16							
				K-40	8.4E-10	1.0E-06						8.4E-18							
				Pu-238	9.9E-12	1.0E-06						9.9E-20							
				Pu-239	2.0E-11	1.0E-06						2.0E-19							
				Ra-226	3.2E-12	1.0E-06						3.2E-20							
				Th-232	2.4E-10	1.0E-06						2.4E-18							
				U-233	2.1E-10	1.0E-06						2.1E-18							
				U-234	1.2E-08	1.0E-06						1.2E-16							
				U-235	5.50E-10	1.0E-06						5.5E-18							
				U-238	1.40E-08	1.0E-06						1.4E-16							
SITE 300 P	OINT SOURCES																		
			ent are conducted on open-air firing ta	bles located at Bun	kers 801 and 851. Th	nese tests ha	ve depleted urani	um material as p	art of the materi	al inventory. There a	are multiple tests p	er year.							
Air activation	on products are cre	eated at the flash x-ray and LINA	AC.																
810A	109	Room Air	Assembly of explosives	U-238	5.4E-02	1.0E-06	NA	NA	NA	None	1	5.4E-08	2360	WSW	2.8E-07	944	SSE	6.7E-06	2
	121		test devices	U-235	6.9E-04	1.0E-06						6.9E-10							
	133			U-234	5.0E-03	1.0E-06						5.0E-09							
	<u> </u>																		
810B	100	Room Air	Assembly of explosives	U-238	1.6E-02	1.0E-06	NA	NA	NA	None	1	1.6E-08	2410	WSW	7.8E-08	907	SSE	2.1E-06	2
			test devices	U-235	2.0E-04	1.0E-06						2.0E-10							
				U-234	1.5E-03	1.0E-06						1.5E-09							
		-																	
			EL 1 1/ (EVD)	N-13	3.4E-03	1.0E+00	NA	NA	NA	None	1	3.4E-03	4114	S	1.8E-08	1809	ENE	3.6E-07	2
801	125	FE-4	Flash X-ray (FXR)		_	1.05.00						2.0E-07							
801	125	FE-4	Flash X-ray (FXR)	Ar-41	2.0E-07	1.0E+00													
801	125	FE-4	Flash X-ray (FXK)	Ar-41	2.0E-07	1.0E+00													
801	125	FE-4	Flash X-ray (FXR)	Ar-41	2.0E-07	1.0E+00													
			Flash X-ray (FXR) ored at the stack. Monitoring data, rat				e emissions.												
*Gross alpha	a and Gross beta e	missions are continuously monit		her than the invent	ory approach, are used	to determin		nd Gross Beta Ra	diation.")										
*Gross alpha	a and Gross beta e	missions are continuously monit	ored at the stack. Monitoring data, rat	her than the invent	ory approach, are used	to determin		nd Gross Beta Ra	diation.") 9.4	НЕРА	0.01	0.0E+00	**	**	0.0E+00	**	**	**	3
*Gross alpha	a and Gross beta e monitoring takes pl	missions are continuously monit lace after HEPA filtration, an una	ored at the stack. Monitoring data, rat bated EDE cannot be determined (see	her than the inventor discussion in Section	ory approach, are used on II, subsection "Stack	to determin	or Gross Alpha ar			HEPA Pre-filter	0.01	0.0E+00 0.0E+00	**	**	0.0E+00	**	**	**	3

***The dose from HTO emis 331 Outside Building 514 is operated by	Table	Stack ID	Operation	Radionuclides	Annual Invantor	Phycical	Stack	Stack	Stack	Control	Control Device	Estimated	10 mrcm/.	Site-Mida Da	se Requirement	01 mram	y Monitoring I	Requirement	Source
Building 331 - As part of D& ***The dose from HTO emis 331 Outside Building 514 is operated by consisting of neutralization,	11	JUNE 10		Radionaciaes	Annual Inventory with Potential for	Physical State	Height (m)	Diameter	Velocity	Device(s)	Abatement	Annual Emissions	Distance to		EDE	Distance	Direction	Unabated	Categor
Building 331 - As part of D& ***The dose from HTO emis 331 Outside Building 514 is operated by consisting of neutralization,	11				Release (Ci)	Factor		(m)	(m/s)	201100(3)	Factor	(Ci)	SWMEI (m)		(mrem)	to MEI (m)	to MEI	EDE (mrem)	Jacogoi
Building 331 - As part of D& ***The dose from HTO emis 331 Outside Building 514 is operated by consisting of neutralization,	11	None	Explosive tests	U-238	1.5E-02	1.0E+00	NA	NA NA	NA NA	None	1	1.5E-02	3170	SSE	1.8E-02	1396	WSW	2.0E-02	4
Building 292 - Diffuse emissi 292 Spill Area Building 331 - As part of D& ***The dose from HTO emis 331 Outside Building 514 is operated by consisting of neutralization,				U-235	2.0E-04	1.0E+00						2.0E-04				-			
Building 292 - Diffuse emissi 292 Spill Area Building 331 - As part of D& ***The dose from HTO emis 331 Outside Building 514 is operated by consisting of neutralization,				U-234	1.4E-03	1.0E+00						1.4E-03							
Building 292 - Diffuse emissi 292 Spill Area Building 331 - As part of D& ***The dose from HTO emis 331 Outside Building 514 is operated by consisting of neutralization,		None	Linear appelenter	N 12	0.25.02	1.05:00	NA	NA	NA	None	1	0 25 02	2170	CCF	1.65.00	2020	FNIF	2 15 00	2
Building 292 - Diffuse emissi 292 Spill Area Building 331 - As part of D& ***The dose from HTO emis 331 Outside Building 514 is operated by consisting of neutralization,		None	Linear accelerator	N-13 O-15	8.2E-02 7.6E-02	1.0E+00 1.0E+00	INA	NA NA	NA	None		8.2E-02 7.6E-02	3170	SSE	1.6E-06	3836	ENE	2.1E-06	2
Building 292 - Diffuse emissi 292 Spill Area Building 331 - As part of D& ***The dose from HTO emis 331 Outside Building 514 is operated by consisting of neutralization,				0-15 Ar-41	7.6E-02 1.5E-04	1.0E+00 1.0E+00	 					7.6E-02 1.5E-04							+
Building 292 - Diffuse emissi 292 Spill Area Building 331 - As part of D& ***The dose from HTO emis 331 Outside Building 514 is operated by consisting of neutralization,				AI-41	1.35-04	1.0E+00	 				 	1.JE-04	1						
Building 292 - Diffuse emissi 292 Spill Area Building 331 - As part of D& ***The dose from HTO emis 331 Outside Building 514 is operated by consisting of neutralization,																			
292 Spill Area Building 331 - As part of D& ***The dose from HTO emis 331 Outside Building 514 is operated by consisting of neutralization,	FUSE SOURCE	ES																	
292 Spill Area Building 331 - As part of D& ***The dose from HTO emis 331 Outside Building 514 is operated by consisting of neutralization,	e emissions ro	secult from tritium-contaminat	ed water which leaked from an under	raround storage tan	Vegetation in the c	rea transniror	water with eleve	ted tritium cons	entrations										
Building 331 - As part of D& ***The dose from HTO emis 331 Outside Building 514 is operated by consisting of neutralization,	e emissions re	esuit iroin tritium-contaminat	su water which leaked from an under	ground Storage tank	vegetation in the ar	ca cranspires	water with elevat	.ea unuum conce	and auOHS.								+	- <u></u> -	
Building 331 - As part of D& ***The dose from HTO emis 331 Outside Building 514 is operated by consisting of neutralization,	Area	None	Evaporation and transpiration	H-3	NA	1	NA	NA	NA	None	1	4.9E-04	1380	ESE	7.2E-08	456	N	1.8E-06	6
***The dose from HTO emis 331 Outside Building 514 is operated by consisting of neutralization,										-						655	W	1.8E-06	
Building 514 is operated by consisting of neutralization,			ment outside the facility is awaiting tr				te Management.												
Building 514 is operated by consisting of neutralization,			IT model; see discussion in Section VI					N. A.	NA	Na		1.05.00	057	ENE	0.75.04	441	CC/4/	2.05.02	+ _
consisting of neutralization,	side	None	Storage of contaminated parts	H-3	NA	1	NA	NA	NA	None	1	1.0E+00	957	ENE	8.7E-04 ***6.5E-04	441	SSW	2.6E-03 ***2.0E-03	6
consisting of neutralization,						 	 								U.JE-U4		-	2.UE-U3	
consisting of neutralization,			l ste Management Division. The waste								ıy involve batch ch	emical treatment							
514 Tank Farm			precipitation, separation, and filtration																
514 Tank Farm																			
	Farm	Area Source	Process liquid hazardous	Am-241	1.5E-05	1.0E-03	NA	NA	NA	None	1	1.5E-08	528	NE	4.1E-04	217	SW	1.2E-03	5
			mixed and radioactive wastes in		4.8E-06	1.0E-03						4.8E-09							
			open topped tanks.	Ba-133 Be-7	3.9E-06 5.0E-07	1.0E-03 1.0E-03	+					3.9E-09 5.0E-10					+		+
				Bi-207	2.6E-07	1.0E-03	 					2.6E-10					 		+
				C-14	6.0E-05	1.0E-03						6.0E-08		+ +			 		+
				Cd-109	2.5E-09	1.0E-03		[2.5E-12							+
				Ce-144	4.0E-05	1.0E-03						4.0E-08							
				Cm-244	4.6E-06	1.0E-03						4.6E-09							
				Co-57	7.8E-07	1.0E-03						7.8E-10							
				Co-58	5.0E-07	1.0E-03				-		5.0E-10				<u> </u>			
				Co-60	3.2E-06	1.0E-03		<u> </u>				3.2E-09							
			+	Cr-51	5.0E-07	1.0E-03						5.0E-10							
			-	Cs-134 Cs-137	4.0E-06 2.5E-05	1.0E-03 1.0E-03						4.0E-09 2.5E-08							-
				Cs-137 Eu-152	2.5E-05 1.5E-05	1.0E-03 1.0E-03	 					2.5E-08 1.5E-08							
				Eu-154	1.5E-05	1.0E-03	 					1.5E-08							
				Eu-155	1.2E-06	1.0E-03						1.2E-09		+ -			 		1
				Fe-55	5.0E-07	1.0E-03						5.0E-10							
				Gd-148	4.0E-06	1.0E-03						4.0E-09							
				H-3	4.5E-04	1.0E-03						4.5E-07							
				Hf-172	5.0E-07	1.0E-03						5.0E-10							
				K-40	2.5E-09	1.0E-03						2.5E-12							
			<u> </u>	Lu-174	5.0E-07	1.0E-03						5.0E-10							
			-	MFP Mp. 54	5.0E-10	1.0E-03						5.0E-13							-
				Mn-54 Na-22	1.2E-06 7.5E-05	1.0E-03 1.0E-03	+					1.2E-09 7.5E-08					+		
				Na-22 Np-237	4.0E-06	1.0E-03	 					7.5E-08 4.0E-09					 		
				P-32	4.1E-03	1.0E-03		[4.1E-06							1
				Pb-210	4.4E-06	1.0E-03						4.4E-09							
				Pu-238	1.0E-05	1.0E-03						1.0E-08							
				Pu-239	4.0E-05	1.0E-03				·		4.0E-08							
			+	Pu-240	1.2E-07	1.0E-03						1.2E-10							+
			+	Pu-241	5.2E-07 4.0E-06	1.0E-03 1.0E-03						5.2E-10 4.0E-09							+
				Pu-242 Pu-244	4.0E-06 1.0E-07	1.0E-03 1.0E-03	+					4.0E-09 1.0E-10		+			+		
				Ra-226	3.2E-07	1.0E-03	 					3.2E-10		+ -			+		
				Sb-125	1.2E-06	1.0E-03						1.2E-09							
				Sc-46	5.0E-07	1.0E-03						5.0E-10							
				Sr-90	6.1E-09	1.0E-03						6.1E-12							
				Tc-99	4.0E-06	1.0E-03	\perp					4.0E-09							
				Th-228	7.0E-10	1.0E-03	<u> </u>	<u> </u>				7.0E-13							
			+	Th-229	4.0E-06	1.0E-03						4.0E-09					 		
			+	Th-230	4.0E-06	1.0E-03						4.0E-09							+
				Th-232 U-232	4.5E+00 4.0E-06	1.0E-03 1.0E-03	+					4.5E-03 4.0E-09		+			+		
		l l		U-232 U-233	4.0E-06 3.9E-06	1.0E-03 1.0E-03	+					4.0E-09 3.9E-09		+ -			+		
				U-234	7.5E-05	1.0E-03	 					7.5E-08					+		
					1.000	1.02.03					+						+		+
					5 5F-06	1.0F-03	1	١.,				5.51-09							
				U-235 U-238	5.5E-06 2.9E-04	1.0E-03 1.0E-03						5.5E-09 2.9E-07							

Property	Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device	Estimated	10 mrem/y S	Site-Wide Do	se Requirement	0.1 mrem	/y Monitoring	Requirement	Source
Description								Height (m)			Device(s)	_					Distance	Direction	Unabated	Category
The description of the ball of a find both Assemble control and control and control of the ball of a find both of the ball of the	The Desilation	C12 Vaud is an au	sand but the Dedicastics and Harr	and and Marke Management Division T	h - Vand i - t f									SWMEI (m)	to SWMEI	(mrem)	to MEI (m)	to MEI	EDE (mrem)	
With the Property of the Pro										outdoors. The d	containers, which are	not air tight, can o	outgas tritium.							
1								e center or the sit	.e.											+
The content of the								NA	NA	NA	None	1	2.3E+00	444	NE	1.1E-02	276	SW	2.4E-02	6
																***8.3E-03			***1.8E-02	
Company of Life Content 1,000 1,																				
Assumbles Accomplished 20-20 (20-20) (19-20) (612	All WAAs*	Area source					NA	NA	NA	None	1		951	ESE	8.5E-09	969	W	5.1E-08	5
Cold 1,250																				
Cold 1233 1662				Accumulation Areas (WAAs)																
C. 66.0 C. 66.1 C. 66.																				
Col. 20 10 10 10 10 10 10 10					Co-57	2.4E-12	1.0E-03													
Color Colo																				
Cart																				
Curist C																				
Birling 1,600 1,																				+
15																				
March Marc																				
March Marc																				
Me53																				
No.229															1					
Post															1					+
Pu-298 1.4-10 1.06-03 1.06-03																				+
Pic-24																				
Pay-26 Ag6-11 106-03					Pu-240	9.5E-12	1.0E-03						9.5E-15							
Representation Repr					Pu-241															
Se-128 3,56-12 1,65-3 1,65-3 1,65-3 1,65-6 1,65-16																				
S-90 1.4C-13 1.0C-03 1.1C-14 1.0C-03 1.1C-14 1.1C-03 1.1C-14																				
To-98																				
19-228 5.9-12 1.06-03 5.9-13 5.9-13																				+
Th-230 5.56-14 1.06-03																				+
Th-212																				+
19-235 2,316-09 1,016-03 2,317-12																				
					U-234	1.7E-08	1.0E-03													
612 Yard Area Source Repackaging operation Am-241 1.9E-08 1.0E-06 NA																				
Ba 138 4.68-10 1.08-06					U-238	1.7E-07	1.0E-03						1.7E-10							
Bar 138 4.68-10 1.06-06	610	VI	A C	Parada di santa di sa	A 241	1.05.00	1.05.00	N/A	N. A.		N		1.05.14	111	NE	1.55.10	20.5	ENE	2.05.10	+
C-14 3.3E-08 1.0E-06 3.3E-14 C-144 7.3E-09 1.0E-06 7.3E-15 C-144 7.3E-09 1.0E-06 7.3E-15 C-1245 5.7E-16 C-245 C-	612	rard	Area Source	Repackaging operation				NA NA	NA NA	NA NA	None	<u>'</u>		444	NE	1.5E-10	295	ENE	2.6E-10	5
Ce-144 7,3E-09 1,0E-06 9,2E-16 9,2E-16																				+
Cm-245 5.7E-10 1.0E-06 5.7E-16 5.7E-16																				1
Co-60					Cm-243	9.2E-10	1.0E-06						9.2E-16							
CS-137 4.4E-09 1.0E-06																				
H3																				
New Year																				
Pu-238 1,3E-09 1,0E-06 1,3E-15																				
Pu-239 9.7E-09 1.0E-06 9.7E-15															-					+
Ra-226															1					+
Th-232 1.6E-09 1.0E-06 1.6E-15																				T
U-234							1.0E-06						1.6E-15							
1.0E-06 1.0E-08 1.0E																				
Comparison Com																				
Comparison Com															1					+
Signature Sign					U-238	4.UE-U8	1.0E-06						4.0E-14							+
Signature Sign	614	Onen Area	Area source	Renackaging of waste	H-3	4 7F-03	1 0F-03	NΔ	NΔ	NΔ	None	1	4 7F-06	420	NNF	2 9F-07	253	ENE	6.5E-07	5
S-35 2.3E-07 1.0E-03 2.3E-10 2.3E-10	317	Open Area	AICA SUUILE					INA	INA	INA	NOUG	1		720	ININE	L.JL-01	233	LINE		+ -
Se-75 2.3E-07 1.0E-03 2.3E-10 Th-232 9.1E-09 1.0E-03 9.1E-12 U-233 4.6E-09 1.0E-03 4.6E-12 Pu-238 2.2E-08 1.0E-03 2.2E-11 Pu-239 1.8E-12 1.0E-03 1.8E-15 Pu-240 1.8E-10 1.0E-03 1.8E-13 Pu-241 4.1E-08 1.0E-03 4.1E-11 Pu-242 9.5E-08 1.0E-03 9.5E-11 U-234 2.3E-09 1.0E-03 2.3E-12 U-235 1.0E-10 1.0E-03 1.0E-13				,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	S-35								2.3E-10							
U-233 4.6E-09 1.0E-03 4.6E-12 Pu-238 2.2E-08 1.0E-03 2.2E-11 Pu-239 1.8E-12 1.0E-03 1.8E-15 Pu-240 1.8E-10 1.0E-03 1.8E-13 Pu-241 4.1E-08 1.0E-03 4.1E-11 Pu-242 9.5E-08 1.0E-03 9.5E-11 U-234 2.3E-09 1.0E-03 2.3E-12 U-235 1.0E-10 1.0E-03 1.0E-13					Se-75	2.3E-07	1.0E-03						2.3E-10							
Pu-238 2.2E-08 1.0E-03 2.2E-11 Pu-239 1.8E-12 1.0E-03 1.8E-15 Pu-240 1.8E-10 1.0E-03 1.8E-13 Pu-241 4.1E-08 1.0E-03 4.1E-11 Pu-242 9.5E-08 1.0E-03 9.5E-11 U-234 2.3E-09 1.0E-03 2.3E-12 U-235 1.0E-10 1.0E-03 1.0E-13					Th-232								9.1E-12							
Pu-239 1.8E-12 1.0E-03 1.8E-15 Pu-240 1.8E-10 1.0E-03 1.8E-13 Pu-241 4.1E-08 1.0E-03 4.1E-11 Pu-242 9.5E-08 1.0E-03 9.5E-11 U-234 2.3E-09 1.0E-03 2.3E-12 U-235 1.0E-10 1.0E-03 1.0E-13																				
Pu-240 1.8E-10 1.0E-03 1.8E-13 Pu-241 4.1E-08 1.0E-03 4.1E-11 Pu-242 9.5E-08 1.0E-03 9.5E-11 U-234 2.3E-09 1.0E-03 2.3E-12 U-235 1.0E-10 1.0E-03 1.0E-13																				
Pu-241 4.1E-08 1.0E-03 4.1E-11 Pu-242 9.5E-08 1.0E-03 9.5E-11 U-234 2.3E-09 1.0E-03 2.3E-12 U-235 1.0E-10 1.0E-03 1.0E-13															1					+
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U-235 1.0E-10 1.0E-03 1.0E-13 1.0E-13																				†
					U-235	1.0E-10	1.0E-03						1.0E-13							
U-238 2.2E-09 1.0E-03 2.2E-12 2.2E-12					U-238	2.2E-09	1.0E-03						2.2E-12							

Building	Room/Area	Stack ID	Operation	Radionuclides	Annual Inventory	Physical	Stack	Stack	Stack	Control	Control Device				se Requirement		/y Monitoring		Source
					with Potential for	State	Height (m)	Diameter	Velocity	Device(s)	Abatement	Annual Emissions	Distance to		EDE	Distance	Direction	Unabated	Categor
					Release (Ci)	Factor		(m)	(m/s)		Factor	(Ci)	SWMEI (m)	to SWMEI	(mrem)	to MEI (m)	to MEI	EDE (mrem)	
The Southea	ast Quadrant of the I	Livermore Site has slightly e	elevated levels of Pu-239 in the surface s	 soil and air (presumal	l oly from resuspension	l n). The source	 e of the Pu-239 v	 vas past waste m	l nanagement opera	ations.									
C. Harris O		A C	D	D 220	N/A	NA	114	NA		None	1	N.A.	0		4.75.04	NIA		NA	
Southeast Q	uadrant	Area Source	Resuspension	Pu-239	NA NA	NA	NA	NA	NA	None	1	NA	0	NA	4.7E-04	NA	NA	NA	6
SITE 300 DI	IFFUSE SOURCES																		
Diffuse sour	ces consist of resus	pension of depleted uraniun	n and waste handling.																
C:+- 200	All	A C	Call resumentian	11.220	NA	NIA	NIA	NA NA	NA	Nama	1	NA	NA	NIA	2.25.02	NA	NIA	NA	
Site 300	All	Area Source	Soil resuspension	U-238 U-235	NA NA	NA NA	NA	NA NA	NA	None	ı	NA NA	NA	NA	3.3E-03	NA	NA	NA	6
			+	U-234	NA NA	NA NA						NA NA							
				0-234	INA	INA						INA INA							
804	Open Area	Area Source	Low-level waste staging area	H-3	NA	NA	NA	NA	NA	None	1	3.9E-04	4508	S	2.1E-07	828	N	4.5E-06	6
				U-238	NA	NA						5.1E-08							
				U-235	NA	NA						6.5E-10							
				U-234	NA	NA						3.1E-09							
EMISSION SO	OURCES THAT ACCO	OUNT FOR MORE THAN 909	6 OF THE POTENTIAL EFFECTIVE DOSE	EQUIVALENT AT EAC	H SITE.														
LIVERMORE	SITE SOURCES																		
612	Yard	Area Source	Storage of low level waste	H-3	NA	NA	NA	NA	NA	None	1	2.3E+00	444	NE	1.1E-02	276	SW	2.4E-02	6
331	All**	Stack 1	Tritium research and development	H-3	*	1	30	1.22	7.59	None	1	2.5E-02	957	ENE	8.1E-03	957	ENE	8.1E-03	3
331	All	Stack 2	Decontamination of parts	H-3	*	1	30	1.22	10.5	None	1	3.6E+01	337	LINL	***5.6E-03	331	LINE	***5.6E-03	
		JUNEAU Z	Decontainination of parts	113			30	1.22	10.5	None	· ·	3.0E+01			3.02 03			3.02.03	
514	Evaporator	Room Air	Waste consolidation	Various nuclides	4.5E-05	1.0E-03	NA	NA	NA	None	1	4.5E-08	528	NE	1.2E-03	217	SW	3.1E-03	1
612	102	Room Air	Laboratory analysis	Various nuclides	4.2E-05	1.0E-03	NA	NA	NA	None	1	4.2E-08	444	NE	1.1E-03	295	ENE	1.5E-03	1
331	Outside	None	Storage of contaminated parts	H-3	NA	1	NA	NA	NA	None	1	1	957	ENE	8.7E-04 ***6.5E-04	441	SSW	0.0026 ***2.0E-03	6
SITE 300 SC	OLIDOTC																		
SITE 300 SC	JURGES																		
851	Firing Table	None	Explosive tests	U-238	1.5E-02	1	NA	NA	NA	None	1	1.5E-02	3170	SSE	1.8E-02	1396	WSW	2.0E-02	4
				U-235	2.0E-04	1	12.7	1	1		·	3.1E-04							<u> </u>
				U-234	1.4E-03	1						2.3E-03							
		Area Source	Soil resuspension	U-238	NA NA	NA	NA	NA	NA	None	1	NA	NA	NA	3.3E-03	NA	NA	NA	6
Sito 200	A II		1 3011 [#5050#[15101]	U-230	INA	INA	I INA	INA	INA	none	1		INA	INA	3.3E-U3	INA	INA	I INA	+ 6
Site 300	All	Area Source		11-225	NA	NIA						NIA						l	
Site 300	All	Alea Jource		U-235 U-234	NA NA	NA NA						NA NA							

ATTACHMENT 2. Surrogate Radionuclides List

The need for selection of a surrogate isotope occurs when an isotope used in operations (isotope of interest) is not contained in the limited nuclide library in the NESHAPs dose compliance model CAP88-PC. The selection of a suitable surrogate is based upon several criteria. If possible, a surrogate isotope is chosen from the CAP88-PC radionuclide library that has a metabolically similar behavior to the isotope of interest. Following an acute inhalation exposure, the metabolically similar surrogate would concentrate in the same specific organs and tissues as the isotope of interest. In most cases the surrogate selected possesses similar modes of decay and decay energies of the radiation type of the isotope of interest. Thus, the surrogate models the behavior of the isotope with similar relative biological effect due to deposition energy.

According to present knowledge, the daughter nuclides produced following physical decay are assumed to remain organ site specific and follow the translocation pathway of the parent. Therefore, when a surrogate of similar metabolic behavior is not available or has a greatly dissimilar half-life, the surrogate chosen is a daughter nuclide of the isotope of interest that will remain organ site specific and follow the translocation pathway of the parent.

Once a surrogate has been selected, the equivalent source term is adjusted by the product of the initial inventory of the isotope of interest and the ratio of the effective dose equivalent of the surrogate to that of the isotope of interest. For determining the dose ratio, the primary exposure pathway is assumed to be that of inhalation and inhalation dose conversion factors (International Commission on Radiological Protection Publication No. 71, "Age-dependent Doses to Members of the Public from Intake of Radionuclides: Part 4 Inhalation Dose Coefficients," Elsevier Science Ltd., 1996) are used for determination of the effective dose equivalents.

In addition, isotopic analysis of mixtures of radionuclides are not always available, and radionuclide usage inventories are stated as "gross alpha," "gross beta," "gross gamma," or "mixed fission products" (MFP). In these cases, ²³⁹Pu is used as the surrogate for gross alpha, ¹³⁷Cs is used as the surrogate for gross gamma, and ⁹⁰Sr is used as the surrogate for gross beta and mixed fission products to provide conservative dose estimates.

Table 2-1 provides a list of radionuclides not in the CAP88-PC library and their respective surrogates.

Table 2-1. List of surrogate radionuclides.

Isotope	Half- Life	Lung Class ^a	ALI (inh) μCi	DAC (inh) μCi/m ³	Surrogate	Half- e Life	Lung Class ^a	ALI (inh) μCi	DAC (inh) μCi/m ³
Ca-108m	127 y	Y	2.0 🗆 101	1.0 🛘 10-8	Co-60	5.271 y	Y	3.0 🗆 101	1.0 🗆 10-8
Bi-207	38 y	W	4.0 □ 102	$1.0 \square 10^{-7}$	Bi-214	19.9 min	W	9.0 □ 102	4.0 □ 10-7
Ca-45	163 d	W	8.0 🛘 102	4.0 □ 10-7	Sr-90	29.12 y	D	2.0 🗆 101	8.0 □ 10 ⁻⁹
Cd-109	464 d	Y	1.0 🛘 102	5.0 🛘 10-8	Co-60	5.271 y	Y	3.0 🛘 101	1.0 □ 10-8
Cf-249	350.6 y	Y	1.0 🛘 10-2	4.0 🗆 10-12	Cm-245	8500 y	W	6.0 🛘 10-3	3.0 🗆 10-12
Cf-250	13.1 y	W	9.0 🛘 10-3	4.0 🗆 10-12	Am-241	432.2 y	W	6.0 🛘 10-3	3.0 🗆 10-12
C1-36	3.01□ 10 ⁵ :	y W	2.0 🛘 102	1.0 🗆 10-7	Cs-137	30 y	D	2.0 🛘 102	6.0 □ 10-8
Es-254	275.7 d	W	7.0 🛘 10-2	3.0 🗆 10-11	Pu-239	24065 y	Y	2.0 🛘 10-2	7.0 🗆 10-12
Eu-149	93.1 d	W	$3.0 \square 10^3$	1.0 🛘 10-6	Pm-151	28.4 hr	Y	$3.0 \square 10^3$	1.0 🗆 10-6
Gd-148	93 y	D	8.0 🛘 10-3	3.0 🗆 10-12	La-140	40.272 h	W	$1.0 \square 10^3$	5.0 🛘 10-7
Os-185	94 d	D	5.0 🛘 102	2.0 🗆 10-7	Mo-99	66 h	Y	$1.0 \square 10^3$	6.0 □ 10-7
P-33	25.4 d	W	$3.0 \square 10^3$	1.0 🛘 10-6	P-32	14.29 d	D	9.0 □ 102	4.0 □ 10-7
Re-184	38 d	W	$1.0 \square 10^3$	6.0 □ 10-7	Mo-99	66 h	Y	$1.0 \square 10^3$	6.0 □ 10-7
Se-75	119.8 d	W	$6.0 \square 10^2$	3.0 🗆 10-7	As-76	26.32 h	W	$1.0 \square 10^3$	6.0 □ 10-7
Sr-85	64.8 d	D	$3.0 \square 10^3$	1.0 □ 10-6	Sr-90	29.12 y	D	$2.0 \square 10^1$	8.0 □ 10 ⁻⁹
Ta-182	115 d	Y	$1.0 \square 10^2$	6.0 □ 10-8	Hf-181	42.4 d	W	$4.0 \square 10^2$	2.0 🗆 10-7
Tb-157	110 y	W	3.0 🛘 102	1.0 🛘 10-7	La-140	40.272 h	W	$1.0 \square 10^3$	5.0 🗆 10-7
Tb-158	180 y	W	$2.0 \square 10^1$	8.0 □ 10 ⁻⁹	La-140	40.272 h	W	$1.0 \square 10^3$	5.0 □ 10 ⁻⁷
T1-204	3.78 y	D	$2.0 \square 10^3$	9.0 🛘 10-7	Pb-214	26.8 min	D	8.0 🛘 102	3.0 □ 10-7
Tm-168	93.1 d	W	$2.0 \square 10^3$	8.0 🗆 10-7	La-140	40.272 h	W	$1.0 \square 10^3$	5.0 □ 10 ⁻⁷
Tm-171	1.92 y	Y	3.0 🛘 102	1.0 🛘 10-7	La-140	40.272 h	W	$1.0 \square 10^3$	5.0 □ 10 ⁻⁷
Y-88	106.64 d	Y	2.0 🛘 102	1.0 🛘 10-7	Y-90	64 h	Y	6.0 □ 102	3.0 □ 10-7
Am-244	10.1 h	W	2.0 🛘 102	8.0 🗆 10-8	Cm-244	18.11 y	W	1.0 🛘 10-2	5.0 🗆 10-12
Au-195	183 d	Y	4.0 □ 102	2.0 🗆 10-7	Ba-133	10.74 y	D	$7.0 \square 10^2$	3.0 □ 10-7
Co-56	78.76 d	Y	2.0 🛘 102	8.0 🗆 10-8	Co-60	5.271 y	Y	3.0 🛘 101	1.0 □ 10-8
Gd-146	48.3 d	W	3.0 🛘 102	1.0 🛘 10-7	Sm-147 1	.06□10 ¹¹ y	v W	4.0 🛘 10-2	2.0 🗆 10-11
Kr-85	10.72 y	Gas	See Note	$1.0 \square 10^{-4}$					
Rh-102	2.9 y	Y	$6.0 \square 10^1$	2.0 🗆 10-8	Rh-106m	29.9 s	Y	$4.0 \square 10^4$	1.0 □ 10-5
U-239	23.54 min	ı Y	$2.0 \square 10^5$	6.0 □ 10 ⁻⁵	U-240	14.1 h	Y	$2.0 \square 10^3$	1.0 🛘 10-6
Zr-90	809 ms	W	N/A	N/A	Y-90	64 h	Y	6.0 🛘 102	3.0 🗆 10-7
Po-209 b	102 y	N/A	N/A	N/A	Pu-239	24065 y	Y	2.0 🛘 10-2	7.0 🗆 10-12

Note: The DAC for Kr-85 also has been relaxed considerably since its beta emission only irradiates the skin. The DAC is based on limitation of non-stochastic effects in the skin; the MPC was derived assuming that the beta particles of energy greater than 0.1 MeV contributed to the whole body dose.

a D = days, W = weeks, Y = years.

b No ALI or DAC information available. Pu-239 used to provide a conservative alpha-emitter dose. Source: Limiting Values of Radionuclide Intake and Air Concentration and Dose Conversion Factors for Inhalation, Submersion and Ingestion, Federal Guidance Report No. 11, EPA-520/1-88-020, U.S. [Invironmental Protection Agency, 1988.

ATTACHMENT 3. Content and Outcome of Proposal to EPA for Use of Air Monitoring in Demonstrating NESHAPs Compliance for Minor Sources



ENVIRONMENTAL PROTECTION DEPARTMENT Operations and Regulatory Affairs Division

March 5, 2003

Jack Broadbent, Director Air Division U.S. Environmental Protection Agency, Region IX 75 Hawthorne St. San Francisco, CA 94105

Subject: Request for Authorization to Use Surveillance Monitoring to Demonstrate Radionuclide NESHAPs Compliance for Minor

Emissions Points

Dear Mr. Broadbent:

Lawrence Livermore National Laboratory (LLNL) is a Department of Energy (DOE) research facility operated by University of California that is subject to requirements of the National Emission Standards for Hazardous Air Pollutants (NESHAPs) of the Clean Air Act, 40 Code of Federal Regulations (CFR) Part 61, Subpart H. LLNL currently demonstrates compliance for various categories of emission sources. These sources include major (continuously monitored) and minor point sources, (i.e., stacks), releases from explosives experiments conducted on open air firing tables, and diffuse area sources, such as contaminated soils. LLNL currently complies with NESHAPs annual dose calculation and reporting requirements through dose estimates based on (1) stack monitoring data for major (continuously monitored) point sources, inventory data/engineering calculations for minor emission point sources, (3) inventory data and empirical scaling laws defining the size and height of explosive tests, and (4) ambient air monitoring or inventory data/engineering calculations for diffuse sources. By this letter, we are requesting authorization to use ambient air surveillance monitoring to demonstrate compliance with the dose assessment requirements of NESHAPs for the source estimates based on inventory data/engineering calculations, i.e., minor emission point sources, and diffuse sources not currently estimated from ambient air.

Several factors justify using air sampler measurements as an alternative method to demonstrate radionuclide NESHAPs compliance.

First, the NESHAPs regulation expressly grants EPA authority to permit compliance demonstration by environmental monitoring. 40 CFR 61.93(b)(5) allows the use of air samplers to obtain "environmental measurements of radionuclide air concentrations at critical receptor locations as an alternative to air dispersion calculations in demonstrating compliance" when the six criteria are established. The program we are proposing will meet the six criteria, which, along with LLNL's plan for meeting them, are submitted as an attachment to this letter. Moreover, the 1995 Memorandum of

TAMM03-021



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Mr. Jack Broadbent, Director Air Division, U.S. EPA Region IX Page 2

Understanding between the U.S. EPA and the U.S. DOE¹ concerning NESHAPs expressly states that the use of environmental measurements of radionuclides at critical receptor locations is "particularly appropriate . . . for facilities with minor emission points (of the periodic confirmatory type) and/or diffuse sources as primary contributors to dose."

EPA has, in fact, granted facilities permission to demonstrate compliance with NESHAPs when the conditions in 40 CFR 61.93(b)(5) are met. The opinion allowing certain operations at the Oak Ridge National Laboratory has been documented on the EPA's web page on the Internet. The Fernald Environmental Management Program and the Mound Plant, with EPA concurrence, have also implemented a NESHAPs compliance demonstration program based on ambient air monitoring.

Secondly, monitoring data provide a better starting point for dose estimates. Air samplers can be placed at or near the location where an individual can be exposed, and air samplers provide measurements of the real concentrations at that location. In contrast, modeling results are estimates of the concentration averaged over an area specified by the model. Moreover, it is important to keep in mind that models are validated, i.e., their accuracy determined, by comparison of modeled results with monitoring data. CAP88-PC, the EPA-approved model currently used to demonstrate NESHAPs compliance, was verified by comparing the environmental monitoring data at five sites with the model predictions. In net effect, the doses calculated for NESHAPs compliance provide a retrospective look at the actual effects of a facility. Monitoring data from continuous ambient air monitors are an excellent source of information about the actual concentrations of radionuclides in air. In fact, LLNL regularly includes in its annual NESHAPs reports a comparison of modeling and monitoring results for the principal emitted radionuclide, tritium, and the comparison shows that model results generally over predict air concentrations at the site perimeter.

Finally, LLNL has collected and measured very low levels of specific nuclides in the ambient air since 1971. Air samplers are currently in use to evaluate diffuse radionuclide emission sources at LLNL for which inventory data is unavailable. It is worthy of note that, for the years 2000 and 2001, diffuse sources (rather than continuously monitored major point sources) have been major contributors to dose at the LLNL Livermore site, accounting for more than one-half of the total dose calculated for the site, and that 40% or more of the total dose calculated for the Livermore site for those two years was based on ambient air measurements.

¹ Memorandum of Understanding Between the U.S. Environmental Protection Agency and the U.S. Department of Energy concerning the Clean Air Act Emission Standards for Radionuclides 40 CFR Part 61 including subparts H, I, Q & T. Signed by the Environmental Protection Agency, September 29, 1994, and by the Department of Energy, April 5, 1995.

² Memorandum, Frank Marcinowski Division Director, Radiation Protection Division, Office of Radiation and Indoor Air, Environmental Protection Agency to Regional Radionuclide NESHAPs Coordinators, Regions I-X, "Criteria to Determine Whether a Leased Facility At DOE is Subject to Subpart H," January 26, 2001 (Found at Applicability Determination Index, Determination Detail Control Number Z010004, http://esdev.sdc-moses.com/oeca/oc/adi/html/Z010004.htm). TAMM03-021

Mr. Jack Broadbent, Director Air Division, U.S. EPA Region IX

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40 CFR 61.93(b)(5) allows the use of air samplers to obtain "environmental measurements of radionuclide air concentrations at critical receptor locations as an alternative to air dispersion calculations in demonstrating compliance" when criteria are established. These criteria, a summary of how they will be met by LLNL, supporting LLNL procedures, as paper copy of the air surveillance monitoring chapter of the LLNL Environmental Report 2001, and a compact disk of the entire LLNL Environmental Report 2001 are submitted with this letter.

LLNL has demonstrated compliance with radionuclide NESHAPs since 1990. At all times, the doses from LLNL operations have been well below the 10 mrem standard. For the Livermore site, the doses have ranged from a high of 0.240 mrem in 1990 to a low of 0.017 mrem reported for calendar year 2001. For Site 300, the doses have ranged from a high of 0.081 mrem in 1994 to a low of 0.019 mrem in 2000. Approval of this application will allow LLNL to make stack monitoring of sources with a potential to emit greater than 10% of the standard the primary focus of its NESHAPS compliance efforts, rather than the current focus on collecting inventory data and modeling nearly 200 sources that account for less than 1% of the total dose consequences from LLNL operations.

We look forward to discussing with you in more detail how our existing monitoring program meets the requirements of 40 CFR 61.93(b)(5) for demonstrating compliance with NESHAPs for minor point sources. Please contact Art Biermann, 925 422-8017 for further information.

Sincerely,

C. Susi Jackson. Leader

Operations and Regulatory Affairs Division

Attachments:

Six Criteria for Use of Environmental Measurements

Air Tritium Sampling Procedure

Air Particulate Sampling Procedure

Air Particulate Sampler Calibration Procedure

Air Surveillance Monitoring Chapter (SAER 2001)

Ambient Air Monitoring Chapter (SAER 2001)

SAER 2001 CD

cc:

Biermann, A.	L-629
Gallegos, G.	L-629
Harrach, B.	L-629
Lessler, R.	EPA IX
Mishra, V.	DOE
Raber, E.	L-626
Rauhut, K.	L-701
Tripodes, J.	L-626
DCC	

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Six Criteria for the Use of Environmental Measurements

40 CFR 61.93(b)(5) allows the use of high-volume air samplers to obtain "environmental measurements of radionuclide air concentrations at critical receptor locations as an alternative to air dispersion calculations in demonstrating compliance" when the following six criteria are established:

- The air at the point of measurement [i.e., the location of the critical receptor] shall be continuously sampled for collection of radionuclides.
- (2) Those radionuclides released from the facility that are the major contributors to the effective dose equivalent must be collected and measured as part of the environmental measurement program.
- (3) Radionuclide concentrations that would cause an effective dose equivalent of 10% of the standard shall be readily detectable and distinguishable from background.
- (4) Net measured radionuclide concentrations shall be compared to the concentration levels in Table 2 appendix E of this part to determine compliance with the standard. In the case of multiple radionuclides being released from a facility, compliance shall be demonstrated if the value for all radionuclides is less than the concentration level in Table 2 of appendix E of this part, and the sum of the fractions that result when each measured concentration value is divided by the value in Table 2 of appendix E of this part for each radionuclide is less than 1.
- (5) A quality assurance program shall be conducted that meets the performance requirements described in appendix B, Method 114 of this part.
- (6) Use of environmental measurements to demonstrate compliance with the standard is subject to prior approval of EPA. Applications for approval shall include a detailed description of the sampling and analytical methodology and show how the above criteria will be met.

Summary of LLNL's ability to meet the six criteria

(1) Continuous sampling

LLNL currently maintains continuous samplers at perimeter and more distant locations, including the location of the site-wide maximally exposed individuals. If the location of the site-wide maximally exposed individual changes, due to a change in operations, or the addition of houses or businesses, samplers can be placed at or near the location, or at a fence line location that reasonably (yet conservatively) measures the concentrations at that location of potential exposure.

(2) Major contributors to dose

LLNL collects samples for tritium and plutonium at the Livermore site, and for tritium and uranium at Site 300. The primary radionuclide emitted from point and diffuse sources at the Livermore site is tritium. Plutonium samples are collected to measure the contribution of the resuspension of contaminated soil in the southeast quadrant of the site. Uranium isotopes are not major contributors to dose at the Livermore site; however, they are the isotopes responsible for more than 90% of the calculated dose for Site 300. Therefore, LLNL maintains a number of high-volume air-particulate samplers at Site 300.

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(3) Radionuclide detection

LLNL maintains a very sensitive air monitoring capability. No single nuclide contributes a dose that is more than 10% of the dose standard. Currently, doses from all nuclides combined are less than 1% of the standard. Detection limits for the analytical methods currently in place for tritium, plutonium and uranium are sufficiently sensitive to provide detections at this level. (Although uranium isotopes are not necessarily distinguishable from background, depending on the filter media, and analytical method, LLNL has recently reinstituted the collection of samples for uranium on cellulose filters, with analysis by mass spectrometry.)

(4) Radionuclide concentrations

Radionuclide concentrations are far below the concentration levels of Table 2 appendix E, and, therefore, well below the NESHAPs standard. According to the "Background Information Document: Procedures for Demonstrating Compliance with 40 CFR Part 61, Subpart I," the values in the table are "the maximum ground-level air concentration that would not result in a dose exceeding the standard." (EPA 520/1-89-001, page 4-28.) In other words, concentrations below the levels of Table 2 would be proportionally lower than the NESHAPs standard. The measured concentrations in calendar year 2001 at the location of the site-wide maximally exposed individual at the Livermore site compared to the Table 2 appendix E concentration levels are provided in Table 1. Note that for Site 300, the maximum monthly concentration of uranium-238 does not take into account the contribution of naturally occurring uranium in resuspended soil, also LLNL is implementing in 2002, a more sensitive method of determining the concentrations of uranium in air based on mass spectrometry rather than alpha spectrometry.

Table 1. Concentrations of nuclides of concern at the location of the maximally exposed included (MEI) in calendar year 2001.

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Location	Nuclide	Table 2 concentration standard	Measured concentration	Measured concentration as a fraction of the standard	Detection limit (approximate)
Livermore site MEI	Tritium	1.5 x 10 ⁻⁹ CVm ³	2.6 x 10 ⁻¹² Ci/m ³	2 x 10 ⁻³	1 x 10 ⁻¹² Ci/m ³
Livermore site MEI	Plutonium- 239		1.2 x 10 ⁻¹⁸ Ci/m ³		3 x 10 ⁻¹⁸ Ci/m ³
Site 300 MEI	Uranium -238	8.3 x 10 ⁻¹⁵ Ci/m ³	2.4 x 10 ⁻¹⁶ Cl/m ³	3 x 10 ⁻²	2 x 10 ⁻¹⁷ Ci/m ³

(5) QA Program

LLNL maintains an extensive QA program. The written procedures, instructions, and the site annual report chapters for calendar year 2001 that address environmental surveillance monitoring are provided as an attachment, and demonstrate compliance with this requirement.

(6) Prior Approval

This lefter and attachments provide a detailed description of the sampling and analytical methodology and show how the preceding five criteria have been, and will be met.

TAMM03-021



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY REGION IX 75 Hawthorne Street San Francisco, CA 94105-3901

April 22, 2003

Ms. C. Susi Jackson, Leader
Operations and Regulatory Affairs Division
Lawrence Livermore National Laboratory
Environmental Protection Department, University of California
P.O. Box 808, Livermore, CA 94551-9900

Subject: Request for Authorization to Use Surveillance Monitoring to Demonstrate Radionuclide NESHAPs Compliance for Minor Emissions Points

Dear Ms. Jackson:

We have reviewed your letter and attachments of March 5, 2003 requesting approval to use surveillance monitoring for minor emission points. In accordance with the provisions of the Clean Air Act and 40 CFR Part 61, Subpart H, your request has been approved.

We request that emissions be closely monitored, identified, and quantified during the use of the approved alternative method, and that the monitoring procedure and related data be kept on file for review by EPA.

This alternative method may be used immediately after this approval is received by the Lawrence Livermore National Laboratory.

If you have any questions, please contact Dick Lessler, at (415) 947-4197

Sincerely,

hick Broadbent Director, Air Division

ach P. Broadley

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